Table 7	7	
ES#	CMCC#	ATCC Deposit#
		Go Bopoolur
85	5175	PTA-1313
86	5176	PTA-1314
87	5177	PTA-1315
88	5178	PTA-1316
89	5179	PTA-1317
90	5180	PTA-1318
91	5181	PTA-1319
92	5182	PTA-1320
93	5183	PTA-1321
94	5184	PTA-1322
95	5185	PTA-1323
96	5186	PTA-1324
97	5187	PTA-1325
98	5188	PTA-1326
99	5189	PTA-1327
100	5190	PTA-1328
101	5191	PTA-1329
102	5192	PTA-1330
103 104	5193 5104	PTA-1331
104	5194 5105	PTA-1332
105	5195 5106	PTA-1333
107	5196 5197	PTA-1334
107	5197	PTA-1335
109	5199	PTA-1336 PTA-1372
110	5200	PTA-1372 PTA-1373
111	5201	PTA-1373
112	5202	PTA-1375
113	5203	PTA-1376
114	5204	PTA-1377
115	5205	PTA-1378
116	5206	PTA-1379
117	5207	PTA-1380
118	5208	PTA-1381
122	5212	PTA-1382
123	5213	PTA-1383
124	5214	PTA-1384
125	5215	PTA-1385
126	5216	PTA-1386
127	5217	PTA-1387
128	5218	PTA-1388
129	5219	PTA-1389
130	5220	PTA-1390
131	5221	PTA-1391
132	5222	PTA-1392
133 134	5223	PTA-1393
134 135	5209 5310	PTA-1431
135 136	5210 5230	PTA-1432
130	5238	PTA-1497

Table 8	Cl. Name	EC No	Clone Nome
ES No.	Clone Name	ES No. ES 109	Clone Name M00027658B:G03
ES 85	M00057077B:D02	ES 109	M00027660C:E03
ES 85	M00057078D:C12		M00027660C:E03
ES 85	M00057079D:E09	ES 109	M00027665B:D01
ES 85	M00057080C:C02	ES 109	
ES 85	M00057085A:A03	ES 109	M00027681D:D02
ES 85	M00057088B:C02	ES 109	M00027699D:D02
ES 85	M00057091A:C03	ES 109	M00027717C:G05
ES 85	M00057091A:C04	ES 109	M00027733D:D05
ES 85	M00057091C:E12	ES 109	M00027742C:B01
ES 85	M00057093B:F09	ES 109	M00027742C:B01
ES 85	M00057099C:C08	ES 109	M00027747D:D01
ES 85	M00057100C:E09	ES 109	M00027757A:B06
ES 85	M00057100D:B03	ES 109	M00027781D:E04
ES 85	M00057103A:E11	ES 109	M00027786D:B01
ES 85	M00057103A:H09	ES 109	M00027803A:H10
ES 85	M00057104B:F08	ES 109	M00027806C:H05
ES 85	M00057106B:A03	ES 109	M00027808D:G10
ES 85	M00057106C:E02	ES 109	M00027817B:B11
ES 85	M00057106D:B06	ES 109	M00027820C:C02
ES 85	M00057108B:F04	ES 109	M00027823C:G07
ES 85	M00057108D:E09	ES 109	M00027829C:D02
ES 85	M00057108D:E09	ES 109	M00027833C:D01
ES 85	M00057112D:B09	ES 110	M00042345A:F12
ES 85	M00057114D:B10	ES 110	M00042523A:C05
ES 85	M00057117D:G11	ES 110	M00042523C:E08
ES 85	M00057118C:C02	ES 110	M00042525D:E01
ES 85	M00057120D:E12	ES 110	M00042527B:D07
ES 85	M00057124B:D10	ES 110	M00042528C:F11
ES 85	M00057127A:F11	ES 110	M00042529C:G07
ES 85	M00057127B:G07	ES 110	M00042532A:F08
ES 85	M00057130C:H11	ES 110	M00042534A:B07
ES 85	M00057131C:B01	ES 110	M00042536D:F01
ES 85	M00057132C:F08	ES 110	M00042537A:H05
ES 85	M00057133D:F01	ES 110	M00042538B:E06
ES 85	M00057134A:C01	ES 110	M00042538D:A08
ES 85	M00057134C:A01	ES 110	M00042539C:E05
ES 85	M00057134D:G10	ES 110	M00042540A:H06
ES 85	M00057135D:H04	ES 110	M00042540D:F03
ES 85	M00057136A:F01	ES 110	M00042540D:H05
ES 85	M00057141B:B02	ES 110	M00042543C:H02
ES 85	M00057141D:D02	ES 110	M00042544B:D02
ES 85	M00057142A:A07	ES 110	M00042544C:F10
ES 85	M00057143C:E05	ES 110	M00042547A:A02
ES 85	M00057145A:D05	ES 110	M00042547B:D11
ES 85	M00057146D:C09	ES 110	M00042547C:F02

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Table 8	CI. NI	EC NI	Clone Name
ES No.	Clone Name	ES No.	Clone Name M00042551A:D09
ES 85	M00057147A:A01	ES 110	
ES 85	M00057150A:C10	ES 110	M00042556A:D04
ES 85	M00057151A:B04	ES 110	M00042563C:E02
ES 86	M00057154A:D06	ES 110	M00042563C:E02
ES 86	M00057154C:B04	ES 110	M00042563D:G09
ES 86	M00057161B:E09	ES 110	M00042564B:H11
ES 86	M00057162A:C07	ES 110	M00042565A:H03
ES 86	M00057162B:H02	ES 110	M00042565C:A08
ES 86	M00057162D:D10	ES 110	M00042567D:C01
ES 86	M00057163D:B01	ES 110	M00042570D:H02
ES 86	M00057165D:E12	ES 110	M00042573C:A07
ES 86	M00057167B:E12	ES 110	M00042574B:H08
ES 86	M00057167B:G12	ES 110	M00042575C:D01
ES 86	M00057167D:B07	ES 110	M00042693D:E04
ES 86	M00057170C:H03	ES 110	M00042694C:E02
ES 86	M00057174B:C06	ES 110	M00042695B:H05
ES 86	M00057174B:G12	ES 110	M00042700B:A01
ES 86	M00057174C:H12	ES 110	M00042700B:D03
ES 86	M00057180A:H11	ES 110	M00042700B:D03
ES 86	M00057181C:D06	ES 110	M00042700D:H05
ES 86	M00057182D:B11	ES 110	M00042704A:F04
ES 86	M00057189B:G05	ES 110	M00042704A:F09
ES 86	M00057191A:A03	ES 110	M00042704D:E02
ES 86	M00057192B:E02	ES 110	M00042705A:D02
ES 86	M00057192D:G02	ES 110	M00042706C:A04
ES 86	M00057196A:E03	ES 110	M00054596B:G11
ES 86	M00057196C:F04	ES 110	M00004101C:H01
ES 86	M00057203C:E06	ES 111	M00042711C:G11
ES 86	M00057208A:A02	ES 111	M00042711D:C04
ES 86	M00057208C:C06	ES 111	M00042712B:B10
ES 86	M00057208C:D08	ES 111	M00042717D:D04
ES 86	M00057211B:F07	ES 111	M00042718B:C03
ES 86	M00057211D:A06	ES 111	M00042720C:D06
ES 86	M00057215B:B02	ES 111	M00042720D:G10
ES 86	M00057217B:B07	ES 111	M00042721A:G07
ES 86	M00057218D:C01	ES 111	M00042727C:H12
ES 86	M00057223C:C06	ES 111	M00042728D:E07
ES 86	M00057224B:C10	ES 111	M00042732A:G09
ES 86	M00057226D:C05	ES 111	M00042735C:G02
ES 86	M00057229D:F06	ES 111	M00042735D:A07
ES 86	M00057230C:D12	ES 111	M00042738B:D10
ES 86	M00057231C:G09	ES 111	M00042739D:D01
	M00037231C.G09 M00057231D:A09	ES 111	M00042741D:D10
ES 86	M00057231D:A09	ES 111	M00042741B:B10
ES 86		ES 111	M00042742C:A06
ES 86	M00057233A:F07	E9 111	19100042742C.A00

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Table 8 ES No.	Clara Nama	EC NI.	Clana Nama
ES 86	Clone Name	ES No. ES 111	Clone Name
ES 86	M00057233B:E04 M00057236B:H06	ES 111	M00042742D:D05 M00042746B:F02
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ES 86	M00057237A:B11	ES 111	M00042746D:B09
ES 86	M00057239A:G08	ES 111	M00042750D:B09
ES 86	M00057241B:B04	ES 111	M00042881D:C08
ES 86	M00057242B:F07	ES 111	M00042883A:F12
ES 87	M00057242D:B09	ES 111	M00042886C:C03
ES 87	M00057242D:H05	ES 111	M00042886C:F01
ES 87	M00057249A:C06	ES 111	M00042887C:D07
ES 87	M00057259A:H10	ES 111	M00042889B:A09
ES 87	M00057259B:B08	ES 111	M00042890D:C08
ES 87	M00057266C:D04	ES 111	M00042891B:C04
ES 87	M00057266C:G12	ES 111	M00042893B:C08
ES 87	M00057268C:E10	ES 111	M00042900C:C07
ES 87	M00057270B:H09	ES 111	M00042901B:A03
ES 87	M00057270C:E04	ES 111	M00042902A:C04
ES 87	M00057271C:E01	ES 111	M00042905A:F11
ES 87	M00057272A:B03	ES 111	M00042905C:C10
ES 87	M00057272C:H04	ES 111	M00042908D:G01
ES 87	M00057272D:A01	ES 111	M00042909B:G04
ES 87	M00057275B:A12	ES 111	M00042911A:H03
ES 87	M00057277B:C09	ES 111	M00042914D:B10
ES 87	M00057277B:E10	ES 111	M00054792D:E09
ES 87	M00057279A:G02	ES 111	M00054793D:B07
ES 87	M00057280C:A06	ES 111	M00054798D:F01
ES 87	M00057283A:E06	ES 111	M00054913C:G03
ES 87	M00057288D:E08	ES 111	M00054915D:E07
ES 87	M00057291C:B06	ES 111	M00054917B:F09
ES 87	M00057297A:F03	ES 111	M00054917D:D12
ES 87	M00057300B:F02	ES 111	M00054918C:D03
ES 87	M00057301B:H12	ES 112	M00054918D:C11
ES 87	M00057304A:E01	ES 112	M00055426B:B02
ES 87	M00057306B:H07	ES 112	M00055426C:H06
ES 87	M00057312B:E11	ES 112	M00055427A:F01
ES 87	M00057318B:B09	ES 112	M00055428C:A02
ES 87	M00057318C:A03	ES 112	M00055429A:H05
ES 87	M00057324A:D12	ES 112	M00055430B:H02
ES 87	M00057325C:C10	ES 112	M00055431C:E09
ES 87	M00057333A:F09	ES 112	M00055438C:C06
ES 87	M00057334B:F01	ES 112	M00055438C:H10
ES 87	M00057337B:G02	ES 112	M00055441B:D02
ES 87	M00057340B:C12	ES 112	M00055445D:G06
ES 87	M00042355A:G02	ES 112	M00055446C:B06
ES 87	M00042355D:C01	ES 112	M00055447D:H04
ES 87	M00042442D:A02	ES 112	M00055447D:H04
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Table 8		l <sub>EG.M</sub>	C1 N
ES No.	Clone Name	ES No.	Clone Name
ES 87	M00042444D:G05	ES 112	M00055448A:D08
ES 87	M00042444D:H08	ES 112	M00055448C:E07
ES 87	M00042450D:H10	ES 112	M00055450A:G09
ES 87	M00042453C:E01	ES 112	M00055450D:B08
ES 87	M00042460D:A07	ES 112	M00055451A:F07
ES 87	M00042517C:F07	ES 112	M00055451A:F11
ES 87	M00042518D:A06	ES 112	M00055451C:G11
ES 87	M00042520A:F04	ES 112	M00055453C:E01
ES 88	M00042520A:F09	ES 112	M00055453C:E01
ES 88	M00042520A:F09	ES 112	M00055454A:A07
ES 88	M00043296C:B10	ES 112	M00055454A:H11
ES 88	M00043300A:H11	ES 112	M00055454C:G05
ES 88	M00043301A:F06	ES 112	M00055456D:F12
ES 88	M00043301D:H09	ES 112	M00055463D:H10
ES 88	M00043304A:D01	ES 112	M00055464A:F05
ES 88	M00043304B:C05	ES 112	M00055466D:B08
ES 88	M00043304B:C05	ES 112	M00055470B:G01
ES 88	M00043306D:B07	ES 112	M00055491A:G08
ES 88	M00043309B:H07	ES 112	M00055494D:C09
ES 88	M00043310C:B03	ES 112	M00055495A:G02
ES 88	M00043313A:A03	ES 112	M00055495C:D05
ES 88	M00043313A:G07	ES 112	M00055495C:F03
ES 88	M00043313D:C06	ES 112	M00055495D:E02
ES 88	M00043314C:H04	ES 112	M00055496A:F09
ES 88	M00043317A:H01	ES 112	M00055496B:E07
ES 88	M00043317C:F04	ES 112	M00055496C:C09
ES 88	M00043323C:D04	ES 112	M00055498A:H09
ES 88	M00043324D:D04	ES 112	M00055500D:B05
ES 88	M00043327D:H02	ES 112	M00055504C:D08
ES 88	M00043327D:H02	ES 112	M00055505D:A10
ES 88	M00043336B:E08	ES 112	M00055508D:E03
ES 88	M00043338A:B03	ES 112	M00055509C:H09
ES 88	M00043338B:A03	ES 112	M00055510B:B07
ES 88	M00043345B:C03	ES 113	M00055511D:E09
ES 88	M00043347B:G12	ES 113	M00055512C:G06
ES 88	M00043349A:C08	ES 113	M00055512D:D07
ES 88	M00043350B:H06	ES 113	M00055512D:F08
ES 88	M00043350C:H09	ES 113	M00055513C:D06
ES 88	M00043352A:E09	ES 113	M00055514D:H05
ES 88	M00043352D:B05	ES 113	M00055516B:E08
ES 88	M00043354D:C01	ES 113	M00055517B:D03
ES 88	M00043355D:H11	ES 113	M00055519B:C06
ES 88	M00043361D:D05	ES 113	M00055519C:H07
ES 88	M00043365A:C06	ES 113	M00055520C:A06
ES 88	M00043374A:B02	ES 113	M00055522A:E07

Table 8		]	
ES No.	Clone Name	ES No.	Clone Name
ES 88	M00043374B:B06	ES 113	M00055522D:C02
ES 88	M00043377A:C03	ES 113	M00055522D:C02
ES 88	M00043379D:C07	ES 113	M00055523D:C03
ES 88	M00043381B:E10	ES 113	M00055525C:B07
ES 88	M00043386D:A06	ES 113	M00055526D:F09
ES 88	M00043388D:C09	ES 113	M00055527C:E02
ES 88	M00043394D:B06	ES 113	M00055527C:E04
ES 88	M00043397B:B02	ES 113	M00055527D:G11
ES 88	M00043397C:B09	ES 113	M00055528C:F06
ES 88	M00043503C:C08	ES 113	M00055529D:B02
ES 88	M00043503C:E05	ES 113	M00055530D:B07
ES 89	M00043504C:G06	ES 113	M00055532C:G08
ES 89	M00043504D:G08	ES 113	M00055534C:H01
ES 89	M00043506A:H09	ES 113	M00055536C:E06
ES 89	M00043507A:D05	ES 113	M00055536C:F03
ES 89	M00043508A:A08	ES 113	M00055538B:H11
ES 89	M00043508D:C01	ES 113	M00055542C:C01
ES 89	M00054486A:B11	ES 113	M00055542C:F06
ES 89	M00054493A:A10	ES 113	M00055542D:A09
ES 89	M00054494A:E01	ES 113	M00055543A:C05
ES 89	M00054496A:B09	ES 113	M00055543A:C05
ES 89	M00054499B:E11	ES 113	M00055543C:G08
ES 89	M00054499B:E11	ES 113	M00055544A:E04
ES 89	M00054502A:D01	ES 113	M00055544B:B02
ES 89	M00054502C:E02	ES 113	M00055545C:H12
ES 89	M00054507A:C11	ES 113	M00055547D:D10
ES 89	M00054510D:H09	ES 113	M00055547D:E05
ES 89	M00054513A:A12	ES 113	M00055548A:F04
ES 89	M00054518D:D03	ES 113	M00055548C:E12
ES 89	M00054520C:B05	ES 113	M00055548C:E12
ES 89	M00054521D:F04	ES 113	M00055552A:C09
ES 89	M00054522B:H11	ES 113	M00055553B:H04
ES 89	M00054523D:A10	ES 113	M00055553D:C07
ES 89	M00054524D:B02	ES 113	M00055553D:H02
ES 89	M00054534D:D02	ES 113	M00055556C:H09
ES 89	M00054535C:H09	ES 113	M00055560B:B12
ES 89	M00054542C:A08	ES 114	M00055560B:F02
ES 89	M00054551C:G03	ES 114	M00055560C:F06
ES 89	M00054555C:G12	ES 114	M00055563A:A02
ES 89	M00054561D:E06	ES 114	M00055572A:B12
ES 89	M00054563B:C09	ES 114	M00055572C:F03
ES 89	M00054568A:G11	ES 114	M00055575A:D08
ES 89	M00054569A:H07	ES 114	M00055578A:H09
ES 89	M00054571C:C01	ES 114	M00055581D:B01
ES 89	M00054572B:C01	ES 114	M00055582B:E04
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Table 8	Τ	Т	T
ES No.	Clone Name	ES No.	Clone Name
ES 89	M00054575C:C01	ES 114	M00055583B:B04
ES 89	M00054580C:D11	ES 114	M00055583B:H05
ES 89	M00054583A:F05	ES 114	M00055584A:C11
ES 89	M00054587A:F09	ES 114	M00055584D:G06
ES 89	M00054590C:G02	ES 114	M00055586C:F05
ES 89	M00054591C:H07	ES 114	M00055586D:F02
ES 89	M00054595A:B02	ES 114	M00055591C:H01
ES 89	M00054595B:H09	ES 114	M00055592D:A05
ES 89	M00054596B:B07	ES 114	M00055594B:A01
ES 89	M00054690D:G07	ES 114 ES 114	M00055597C:E08
ES 89	M00054600D:G07	ES 114 ES 114	M00055601C:D09
ES 89	M00054601A:H10	ES 114 ES 114	M00055602B:G10
ES 89	M00054602A:C04	ES 114 ES 114	M00055602C:E07
ES 90	M00054602B:D02	ES 114	M00055609A:G03
ES 90	M00054604A:D09	ES 114 ES 114	M00055609D:F12
ES 90	M00054604A:D09	ES 114	M00055613A:D10
ES 90	M00054605C:D01	ES 114	M00055613A:E02
ES 90	M00054609A:F01	ES 114	M00055618C:A06
ES 90	M00054609D:H06	ES 114	M00055628A:A08
ES 90	M00054611C:F02	ES 114	M00055630B:E09
ES 90	M00054613A:D09	ES 114	M00055633D:A02
ES 90	M00054613A:D09	ES 114	M00055633D:G11
ES 90	M00054617B:A09	ES 114	M00055635A:H10
ES 90	M00054621B:C06	ES 114	M00055635B:E10
ES 90	M00054621D:D11	ES 114	M00055635C:G04
ES 90	M00054629C:E09	ES 114	M00055636A:F10
ES 90	M00054636B:B03	ES 114	M00055647C:B04
ES 90	M00054636C:A02	ES 114	M00055653A:H04
ES 90	M00054636C:F02	ES 114	M00055656A:E09
ES 90	M00054638A:D09	ES 114	M00055662C:A04
ES 90	M00054638B:C08	ES 114	M00055664C:A08
ES 90	M00054646C:B01	ES 114	M00055668B:B07
ES 90	M00054647D:H02	ES 114	M00055679A:A07
ES 90	M00054648C:H10	ES 114	M00055681B:G02
ES 90	M00054660D:F05	ES 114	M00055685D:E01
ES 90	M00054665B:H08	ES 114	M00055686D:E11
ES 90	M00054665D:E11	ES 114	M00055687C:F01
ES 90	M00054677C:D02	ES 114	M00055688C:B04
ES 90	M00054678A:E07	ES 114	M00055689A:G12
ES 90	M00054679B:D12	ES 115	M00055689D:F02
ES 90	M00054680B:E06	ES 115	M00055689D:F07
ES 90	M00054680D:B11	ES 115	M00055691B:E07
ES 90	M00054681C:B02	ES 115	M00055692D:E07
ES 90	M00054684C:H12	ES 115	M00055701C:D10
ES 90	M00054689D:E12	ES 115	M00055703A:B08

Table 8			<del></del>
ES No.	Clone Name	ES No.	Clone Name
ES 90	M00054691A:E05	ES 115	M00055703B:B06
ES 90	M00054692B:D01	ES 115	M00055703B:C05
ES 90	M00054694D:G04	ES 115	M00055703C:G09
ES 90	M00054706B:C09	ES 115	M00055704C:D07
ES 90	M00054707B:B08	ES 115	M00055705C:G07
ES 90	M00054707B:E05	ES 115	M00055706A:A01
ES 90	M00054707B.E03	ES 115	M00055706B:G01
ES 90	M00054713A:D12	ES 115	M00055707D:C08
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ES 90	M00054720D:F11	ES 115	M00055709B:G09
ES 90	M00054721C:F11	ES 115	M00055716C:B04
ES 90	M00054722C:D01	ES 115	M00055717B:F04
ES 90	M00054722D:C08	ES 115	M00055718A:F05
ES 90	M00054726A:F08	ES 115	M00055720B:G09
ES 90	M00054727D:E10	ES 115	M00055720C:A06
ES 90	M00054727D:H06	ES 115	M00055720D:A01
ES 90	M00054728B:E08	ES 115	M00055721B:F06
ES 91	M00054728D:B10	ES 115	M00055721B:F06
ES 91	M00054729A:E01	ES 115	M00055721C:E05
ES 91	M00054731C:C12	ES 115	M00055723A:B08
ES 91	M00054732D:E03	ES 115	M00055723D:E05
ES 91	M00054734D:H10	ES 115	M00055724D:D09
ES 91	M00054739A:G03	ES 115	M00055726B:B08
ES 91	M00054739C:D03	ES 115	M00055726C:D12
ES 91	M00054739C:E06	ES 115	M00055726C:G10
ES 91	M00054740A:H08	ES 115	M00055729D:A06
ES 91	M00054741A:C10	ES 115	M00055731A:H12
ES 91	M00054741A:E10	ES 115	M00055733A:G11
ES 91	M00054741D:G10	ES 115	M00055734C:H05
ES 91	M00054743C:E02	ES 115	M00055735C:C07
ES 91	M00054745D:A03	ES 115	M00055735C:G05
ES 91	M00054747A:F01	ES 115	M00055736A:D06
ES 91	M00054747D:C06	ES 115	M00055736B:G03
ES 91	M00054750C:D12	ES 115	M00055736C:G07
ES 91	M00054752B:A07	ES 115	M00055740B:B12
ES 91	M00054755B:H06	ES 115	M00055740B:F09
ES 91	M00054759A:B08	ES 115	M00055743B:C12
ES 91	M00054760A:A12	ES 115	M00055744B:C08
ES 91	M00054762B:F07	ES 115	M00055744C:F08
ES 91	M00054765B:C05	ES 115	M00055744C:F09
ES 91	M00054766C:B04	ES 115	M00055744D:G08
ES 91	M00054769A:F07	ES 115	M00055747C:D09
ES 91	M00054772C:C06	ES 115	M00055749D:H11
ES 91	M00054773A:A12	ES 116	M00055751C:D01
ES 91	M00054776B:F01	ES 116	M00055755C:D01
ES 91	M00054779A:F07	ES 116	M00055755D:H03
71	14400034773A.TU/	153 110	בטוו.עכבו ככטטטייון

Table 0	T	<del></del>	T
Table 8 ES No.	Clone Name	ES No.	Clone Name
ES 91	M00054780C:G08	ES 116	M00055761D:C03
ES 91	M00054781B:B04	ES 116	M00055763B:E06
ES 91	M00054802A:G02	ES 116	M00055765A:B05
ES 91	M00054804D:H12	ES 116	M00055766A:H03
ES 91	M00054808A:D07	ES 116	M00055768A:B05
ES 91	M00054808B:F08	ES 116	M00055770C:G01
ES 91	M00054810B:H02	ES 116	M00055771A:A11
ES 91	M00054810B:A05	ES 116	M00055771A:D01
ES 91	<del></del>	ES 116	M00055771C:A11
<del></del>	M00054812D:C07	ES 116	M00055771C:A11
ES 91	M00054812D:C07		<del></del>
ES 91	M00054815C:E01	ES 116 ES 116	M00055771D:D04
ES 91	M00054816C:D11	<del></del>	M00055771D:F07
ES 91	M00054821A:C11	ES 116	M00055774C:E01
ES 91	M00054823D:H07	ES 116	M00055774D:G03
ES 91	M00054826B:C10	ES 116	M00055775C:B10
ES 91	M00054826B:E05	ES 116	M00055775D:C06
ES 91	M00054826D:C10	ES 116	M00055778A:F09
ES 91	M00054827B:H01	ES 116	M00055779B:A02
ES 92	M00054832D:E09	ES 116	M00055780B:B07
ES 92	M00054836A:B05	ES 116	M00055780D:G08
ES 92	M00054839B:B02	ES 116	M00055781C:C05
ES 92	M00054839C:F06	ES 116	M00055782A:F02
ES 92	M00054841D:B07	ES 116	M00055783A:C11
ES 92	M00054841D:B07	ES 116	M00055785B:F03
ES 92	M00054842D:C11	ES 116	M00055785C:E08
ES 92	M00054844D:F06	ES 116	M00055786A:D05
ES 92	M00054849D:H11	ES 116	M00055788D:A03
ES 92	M00054851B:E03	ES 116	M00055790C:H02
ES 92	M00054854D:E08	ES 116	M00055791A:D05
ES 92	M00054856D:A02	ES 116	M00055791D:F03
ES 92	M00054857D:E12	ES 116	M00055792B:G09
ES 92	M00054862B:B07	ES 116	M00055792D:E07
ES 92	M00054863B:G03	ES 116	M00055794A:G11
ES 92	M00054865B:H04	ES 116	M00055794C:B06
ES 92	M00054866C:G07	ES 116	M00055796D:E10
ES 92	M00054867A:C07	ES 116	M00055797A:D08
ES 92	M00054867B:B02	ES 116	M00055797B:E07
ES 92	M00054867C:B07	ES 116	M00055798B:C06
ES 92	M00054869C:D01	ES 116	M00055800A:D08
ES 92	M00054870B:D09	ES 116	M00055800B:C08
ES 92	M00054875B:C04	ES 116	M00055802A:D08
ES 92	M00054876B:G03	ES 116	M00055802A:G02
ES 92	M00054877A:H12	ES 116	M00055802B:H04
ES 92	M00054895B:D09	ES 116	M00055802C:F12
ES 92	M00054899D:F07	ES 116	M00055803B:A11

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Table 8		200	C1 >1
ES No.	Clone Name	ES No.	Clone Name
ES 92	M00054899D:G01	ES 117	M00055803C:D08
ES 92	M00054903D:C12	ES 117	M00055804A:F03
ES 92	M00054908B:F07	ES 117	M00055804B:F01
ES 92	M00054910D:G06	ES 117	M00055805B:C07
ES 92	M00054926D:F01	ES 117	M00055805C:D10
ES 92	M00054927B:E08	ES 117	M00055806A:H12
ES 92	M00054931C:A09	ES 117	M00055806B:F07
ES 92	M00054933A:D07	ES 117	M00055806C:E09
ES 92	M00054934C:D03	ES 117	M00055807B:G10
ES 92	M00054935A:E01	ES 117	M00055807B:G10
ES 92	M00054935A:G04	ES 117	M00055807D:C04
ES 92	M00054937A:B03	ES 117	M00055808C:G11
ES 92	M00054937B:A12	ES 117	M00055811A:A08
ES 92	M00054937B:F03	ES 117	M00055811D:C12
ES 92	M00054937C:B10	ES 117	M00055812A:E01
ES 92	M00054941C:G04	ES 117	M00055814C:D11
ES 92	M00054943C:C04	ES 117	M00055816B:F01
ES 92	M00054943D:C03	ES 117	M00055817B:F01
ES 92	M00054945C:G07	ES 117	M00055817C:C08
ES 92	M00054947B:G12	ES 117	M00055817C:D08
ES 92	M00054949A:E03	ES 117	M00055818A:F12
ES 93	M00054949C:A07	ES 117	M00055818D:E10
ES 93	M00054950D:G06	ES 117	M00055820A:E08
ES 93	M00054952A:F01	ES 117	M00055820B:E05
ES 93	M00054952C:H06	ES 117	M00055820C:C08
ES 93	M00054953D:G10	ES 117	M00055820D:G10
ES 93	M00054954B:C03	ES 117	M00055821A:A06
ES 93	M00054954D:F01	ES 117	M00055821A:G12
ES 93	M00054957A:B02	ES 117	M00055822B:H04
ES 93	M00054959C:C11	ES 117	M00055823B:D03
ES 93	M00054963C:H11	ES 117	M00055823C:D11
ES 93	M00054963D:H04	ES 117	M00055825B:E03
ES 93	M00054964A:H11	ES 117	M00055826A:F04
ES 93	M00054965B:H02	ES 117	M00055827B:D02
ES 93	M00054970D:G03	ES 117	M00055827D:A02
ES 93	M00054973B:A10	ES 117	M00055827D:C06
ES 93	M00054975C:C04	ES 117	M00055827D:E05
ES 93	M00054980D:C02	ES 117	M00055829C:G09
ES 93	M00054981C:E11	ES 117	M00055830A:G10
ES 93	M00054981D:C06	ES 117	M00055832D:E12
ES 93	M00054984D:B12	ES 117	M00055833D:A11
ES 93	M00054984D:C07	ES 117	M00055838B:D12
ES 93	M00054985C:F07	ES 117	M00055838B:G12
ES 93	M00054987D:C02	ES 117	M00055839A:F09
ES 93	M00054988C:G02	ES 117	M00055839B:A10
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Table 8	Ţ	<del></del>	T
ES No.	Clone Name	ES No.	Clone Name
ES 93	M00054995A:C10	ES 117	M00055840C:D06
ES 93	M00054996C:B11	ES 117	M00055841A:B09
ES 93	M00054996C:C09	ES 117	M00055841C:D05
ES 93	M00054997C:B12	ES 118	M00055848C:H06
ES 93	M00054997C:H03	ES 118	M00055849C:D08
ES 93	M00055000C:F04	ES 118	M00055850B:F03
ES 93	M00055002D:E04	ES 118	M00055851A:G11
ES 93	M00055005B:H11	ES 118	M00055851C:F12
ES 93	M00055005D:B08	ES 118	M00055851C:F12
ES 93	M00055008A:B08	ES 118	M00055852A:A07
ES 93	M00055008D:B09	ES 118	M00055854A:E04
ES 93	M00055011C:E04	ES 118	M00055856A:F04
ES 93	M00055017A:A11	ES 118	M00055856C:F07
ES 93	M00055021D:D11	ES 118	M00055860D:A08
ES 93	M00055022A:H04	ES 118	M00055861A:D03
ES 93	M00055027B:D07	ES 118	M00055864A:E11
ES 93	M00055027D:F08	ES 118	M00055864A:H02
ES 93	M00055032D:A06	ES 118	M00055866D:A02
ES 93	M00055034C:G01	ES 118	M00055868A:F06
ES 93	M00055034D:H01	ES 118	M00055868D:D03
ES 93	M00055037A:E10	ES 118	M00055868D:F09
ES 93 ES 93	M00055039A:G01	ES 118 ES 118	M00055869B:A06 M00055871A:F05
ES 93	M00055039C:E02 M00055041A:E02	ES 118	M00055871D:G06
ES 94	M00055042A:B01	ES 118	M00055871D:000
ES 94	M00055046B:C07	ES 118	M00055873A:B11
ES 94	M00055046C:E11	ES 118	M00055873B:E03
ES 94	M00055050C:G04	ES 118	M00055874B:B06
ES 94	M00055053C:B03	ES 118	M00055874D:D03
ES 94	M00055054A;C02	ES 118	M00055879B:E11
ES 94	M00055056D:B06	ES 118	M00055879C:D04
ES 94	M00055057A:F03	ES 118	M00055880D:F12
ES 94	M00055063D:G01	ES 118	M00055882C:A06
ES 94	M00055064A:E12	ES 118	M00055882C:A09
ES 94	M00055071B:A02	ES 118	M00055884A:E10
ES 94	M00055073C:H12	ES 118	M00055884C:B01
ES 94	M00055075B:H05	ES 118	M00055884D:A05
ES 94	M00055077C:F11	ES 118	M00055886D:G09
ES 94	M00055085A:A10	ES 118	M00055887A:C06
ES 94	M00055087A:A10	ES 118	M00055887A:F07
ES 94	M00055088A:A12	ES 118	M00055887B:E04
ES 94	M00055088C:E09	ES 118	M00055888B:B05
ES 94	M00055093B:H05	ES 118	M00055889C:H12
ES 94	M00055094B:H09	ES 118	M00055891B:A04
ES 94	M00055097A:G06	ES 118	M00055893B:C05

Table 8	<del>,</del>		T
ES No.	Clone Name	ES No.	Clone Name
ES 94	M00055100B:D08	ES 118	M00055896C:F11
ES 94	M00055104C:B12	ES 118	M00055900B:B05
ES 94	M00055106A:D07	ES 118	M00055906B:D10
ES 94	M00055100A.D07	ES 118	M00055906C:F12
ES 94	M00033111B:D03	ES 118	M00055908B:H11
ES 94	M00055113B:A11	ES 118	M00055908C:E12
ES 94	<del></del>	ES 122	M00056209B:F12
<u> </u>	M00055114D:A11		<del></del>
ES 94	M00055115A:E05	ES 122 ES 122	M00056210B:E03 M00056212C:G01
ES 94	M00055116B:B02		<del></del>
ES 94	M00055117C:C03	ES 122	M00056213A:A04
ES 94	M00055121D:C07	ES 122	M00056215A:E03
ES 94	M00055125B:E06	ES 122	M00056218C:G03
ES 94	M00055125B:F01	ES 122	M00056220D:D02
ES 94	M00055128D:B10	ES 122	M00056220D:D09
ES 94	M00055130D:G01	ES 122	M00056221D:E05
ES 94	M00055131C:B10	ES 122	M00056222C:F02
ES 94	M00055134B:E03	ES 122	M00056223B:G03
ES 94	M00055134B:H02	ES 122	M00056224C:B10
ES 94	M00055134D:B03	ES 122	M00056224D:E08
ES 94	M00055137C:C04	ES 122	M00056225D:G09
ES 94	M00055145A:F07	ES 122	M00056226C:F12
ES 94	M00055148D:D11	ES 122	M00056228B:A07
ES 94	M00055154C:F04	ES 122	M00056231B:G09
ES 94	M00055157A:C11	ES 122	M00056231B:G09
ES 94	M00055161D:A11	ES 122	M00056232C:E06
ES 94	M00055162A:F06	ES 122	M00056232D:G12
ES 94	M00055163A:C02	ES 122	M00056233D:F03
ES 95	M00055170A:F01	ES 122	M00056236D:G01
ES 95	M00055170D:E02	ES 122	M00056238B:E07
ES 95	M00055172D:D04	ES 122	M00056243C:G10
ES 95	M00055179C:D02	ES 122	M00056244A:B08
ES 95	M00055181A:E01	ES 122	M00056244B:C07
ES 95	M00055182B:C07	ES 122	M00056246A:B03
ES 95	M00055185C:B01	ES 122	M00056246C:G07
ES 95	M00055194D:C05	ES 122	M00056247A:G06
ES 95	M00055196B:A09	ES 122	M00056248A:A09
ES 95	M00055198D:F07	ES 122	M00056250B:F01
ES 95	M00055198D:G07	ES 122	M00056251C:A06
ES 95	M00055201D:A03	ES 122	M00056252B:H08
ES 95	M00055201D:B07	ES 122	M00056252D:E11
ES 95	M00055203B:H02	ES 122	M00056253A:F12
ES 95	M00055206A:H04	ES 122	M00056253D:H06
ES 95	M00055207D:A04	ES 122	M00056254A:H02
ES 95	M00055209D:A08	ES 122	M00056256C:H11
ES 95	M00055209D:D10	ES 122	M00056262B:B08

Table 8			
ES No.	Clone Name	ES No.	Clone Name
ES 95	M00055216A:A03	ES 122	M00056263D:C01
ES 95	M00055216A:A03	ES 122	M00056267A:E02
ES 95	M00055222D:H05	ES 122	M00056267C:B02
ES 95	M00055227A:H09	ES 122	M00056268B:B04
ES 95	M00055227D:E02	ES 122	M00056268C:D01
ES 95	M00055227D:E07	ES 122	M00056273B:A01
ES 95	M00055231A:D10	ES 122	M00056281D:E04
ES 95	M00055242A:E06	ES 122	M00056282A:F11
ES 95	M00055242B:A01	ES 122	M00056282B:D04
ES 95	M00055242D:D04	ES 123	M00056282D:C01
ES 95	M00055243A:F04	ES 123	M00056282D:H09
ES 95	M00055243A:G01	ES 123	M00056283A:E02
ES 95	M00055245B:A09	ES 123	M00056283A:E02
ES 95	M00055247B:A11	ES 123	M00056283D:C03
ES 95	M00055252A:C02	ES 123	M00056286A:E08
ES 95	M00055259D:F04	ES 123	M00056286D:A12
ES 95	M00055260B:A05	ES 123	M00056290B:F05
ES 95	M00055260C:F12	ES 123	M00056290D:E08
ES 95	M00055262C:B11	ES 123	M00056290D:H02
ES 95	M00055263A:G09	ES 123	M00056291B:G01
ES 95	M00055271D:C05	ES 123	M00056291D:B05
ES 95	M00055273B:C05	ES 123	M00056292B:E05
ES 95	M00055274C:F02	ES 123	M00056293B:E08
ES 95	M00055279B:G08	ES 123	M00056293C:F05
ES 95	M00055279C:E12	ES 123	M00056293C:G09
ES 95	M00055283B:F05	ES 123	M00056295A:F07
ES 95	M00055283C:H02	ES 123	M00056295C:D06
ES 95	M00055289B:D02	ES 123	M00056300A:A05
ES 95	M00055294B:D04	ES 123	M00056302B:F12
ES 96	M00055302D:F02	ES 123	M00056303A:C03
ES 96	M00055306A:G09	ES 123	M00056303C:B04
ES 96	M00055319B:A01	ES 123	M00056304D:G11
ES 96	M00055322B:E01	ES 123	M00056307A:H12
ES 96	M00055324C:H10	ES 123	M00056310B:G06
ES 96	M00055325C:B12	ES 123	M00056312B:A04
ES 96	M00055327D:H08	ES 123	M00056312D:C03
ES 96	M00055332C:G11	ES 123	M00056313C:F07
ES 96 ES 96	M00055334C:H09 M00055335A:H03	ES 123	M00056319C:G01 M00056320C:H02
ES 96	M00055335A:H03 M00055338B:H07	ES 123 ES 123	M00056323A:H10
ES 96	<del></del>	ES 123	M00056323A:H10
ES 96	M00055344C:H09	ES 123	M00056323C:C12
ES 96	M00055345C:H11 M00055346B:D02	ES 123	M00056323C:C12
ES 96	M00055350A:F01	ES 123	M00056324B:D02
ES 96	M00055356C:C06	ES 123	M00056330C:D03
Tro 30	M00033330C.C00	123	M100030330C.D03

Table 8	Т		T
ES No.	Clone Name	ES No.	Clone Name
<u> </u>	M00055358B:C01		M00056331B:D01
ES 96	<del></del>	ES 123	
ES 96	M00055361D:H01	ES 123	M00056338B:B06
ES 96	M00055363D:G12	ES 123	M00056338C:B10
ES 96	M00055364D:E01	ES 123	M00056342A:G05
ES 96	M00055368B:C10	ES 123	M00056342B:G03
ES 96	M00055368C:B06	ES 123	M00056342C:F11
ES 96	M00055371B:F01	ES 123	M00056344A:G03
ES 96	M00055373D:D10	ES 123	M00056345D:E03
ES 96	M00055374A:A08	ES 123	M00056437C:H07
ES 96	M00055376B:B01	ES 123	M00056438C:A06
ES 96	M00055379D:C08	ES 123	M00056447B:A04
ES 96	M00055381B:E09	ES 123	M00056448B:C09
ES 96	M00055383B:B04	ES 124	M00056456C:A09
ES 96	M00055383B:B04	ES 124	M00056456C:F02
ES 96	M00055384D:A03	ES 124	M00056456D:F01
ES 96	M00055385C:G06	ES 124	M00056459A:C07
ES 96	M00055388A:G08	ES 124	M00056459A:C07
ES 96	M00055388A:H08	ES 124	M00056459A:D07
ES 96	M00055390B:D08	ES 124	M00056460A:G11
ES 96	M00055391A:G08	ES 124	M00056466A:A03
ES 96	M00055391C:G06	ES 124	M00056466A:E02
ES 96	M00055395A:C02	ES 124	M00056467C:E07
ES 96	M00055396A:G07	ES 124	M00056475B:C12
ES 96	M00055404D:C07	ES 124	M00056475C:F01
ES 96	M00055405A:D09	ES 124	M00056475C:F02
ES 96	M00055405B:H05	ES 124	M00042432C:H10
ES 96	M00055405D:G07	ES 124	M00042440A:E05
ES 96	M00055406B:D05	ES 124	M00042461A:A10
ES 96	M00055408B:E09	ES 124	M00042463A:F09
ES 96	M00055408D:F03	ES 124	M00042466D:H06
ES 96	M00055413A:B07	ES 124	M00042469D:H04
ES 96	M00055414C:A11	ES 124	M00042511A:A04
ES 97	M00055415B:H11	ES 124	M00042513D:A12
ES 97	M00055417A:G08	ES 124	M00042515C:A10
ES 97	M00055419D:G01	ES 124	M00042756B:B01
ES 97	M00055420A:E06	ES 124	M00042758D:H12
ES 97	M00055420B:F10	ES 124	M00042760B:C07
ES 97	M00055420D:G04	ES 124	M00042764B:B04
ES 97	M00055421B:D04	ES 124	M00042767D:D02
ES 97	M00055421C:C11	ES 124	M00042770B:B12
ES 97	M00055423A:A10	ES 124	M00042771B:A03
ES 97	M00055423A:G08	ES 124	M00042777A:D06
ES 97	M00055423C:H10	ES 124	M00042781C:A06
ES 97	M00055425D:C05	ES 124	M00042783C:A03
ES 97	M00055472A:F02	ES 124	M00042787C:E09
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Table 8	<del></del>		T
ES No.	Clone Name	ES No.	Clone Name
ES 97	M00055472B:H03	ES 124	M00042792C:G10
ES 97	M00055475D:G08	ES 124	M00042792D:F04
ES 97	M00055479A:G02	ES 124	M00042793B:G06
ES 97	M00055479C:C12	ES 124	M00042793B:G00 M00042800A:A03
ES 97	M00055480C:H10	ES 124	M00042801B:B06
ES 97	M00055482D:A01	ES 124	M00042802C:G11
ES 97	M00055484A:G07	ES 124	M00042805A:E06
ES 97	M00055485A:C09	ES 124	M00042805D:H08
ES 97	M00055487B:F06	ES 124	M00042814D:B11
ES 97	M00001340A:E01	ES 124	M00042816B:F04
ES 97	M00001470C:G01	ES 124	M00042818A:E12
ES 97	M00001470C:G01	ES 124	M00042823B:G04
ES 97	M00001491B:C08	ES 124	M00042826B:C05
ES 97	M00001537D:F10	ES 124	M00042826D:C03
ES 97	M00001537D:F10	ES 124	M00042833D:G01
ES 97	M00001561B:G01	ES 125	M00042834C:B06
ES 97	M00001625A:B08	ES 125	M00042835C:C01
ES 97	M00001637A:D09	ES 125	M00042835D:D02
ES 97	M00003792B:A11	ES 125	M00042838D:E11
ES 97	M00003794C:D07	ES 125	M00042842A:C01
ES 97	M00003804D:A09	ES 125	M00042842D:E08
ES 97	M00003922B:H03	ES 125	M00042844C:C12
ES 97	M00003948A:B12	ES 125	M00042846C:D09
ES 97	M00003986D:G12	ES 125	M00042848D:G12
ES 97	M00003986D:G12	ES 125	M00042849D:F11
ES 97	M00004054C:G05	ES 125	M00042850B:C04
ES 97	M00004066C:D02	ES 125	M00042850C:C10
ES 97	M00004080A:A05	ES 125	M00042853A:G03
ES 97	M00004087D:B11	ES 125	M00042853D:A04
ES 97	M00004093D:C10	ES 125	M00042855A:B09
ES 97	M00004167C:D11	ES 125	M00042856C:F07
ES 97	M00004167C:D11	ES 125	M00042864A:E05
ES 97	M00004198B:A11	ES 125	M00042867D:H01
ES 97	M00004296D:G11	ES 125	M00042869C:E06
ES 98	M00004304A:D07	ES 125	M00042875C:E04
ES 98	M00004842C:B07	ES 125	M00042879B:F09
ES 98	M00004850C:G05	ES 125	M00056346C:C12
ES 98	M00004852D:A04	ES 125	M00056351B:D06
ES 98	M00004868B:D12	ES 125	M00056356B:F04
ES 98	M00004869D:D06 M00004971B:G04	ES 125	M00056359C:A11
ES 98 ES 98	M00004971B:G04 M00004972C:E01	ES 125	M00056362D:G05
ES 98	M00004972C:E01 M00005000B:H08	ES 125 ES 125	M00056363A:B06
ES 98	<del></del>	ES 125 ES 125	M00056368C:F04
	M00005019D:D02		M00056369B:D12
ES 98	M00005293B:D06	ES 125	M00056370B:G02

Table 8         ES No.         Clone Name         ES No.         Clone Name           ES 98         M00005309D:E05         ES 125         M00056371B:F12           ES 98         M00005313C:B02         ES 125         M00056374B:H02           ES 98         M000053373D:H06         ES 125         M0000563874B:H02           ES 98         M000053385C:A10         ES 125         M00056383A:C10           ES 98         M00005389C:C11         ES 125         M00056410B:E04           ES 98         M00005395A:D09         ES 125         M00056410B:E04           ES 98         M00005395A:D09         ES 125         M0005641CC:B12           ES 98         M00005406C:A11         ES 125         M0005641CC:B12           ES 98         M00005407B:E12         ES 125         M0005642CD:D07           ES 98         M00005417C:C07         ES 125         M0005642A:F12           ES 98         M00005413B:F03         ES 125         M0005642A:F12           ES 98         M00005412C:F12         ES 125         M0005642A:F12           ES 98         M00005449C:C10         ES 125         M0005642B:H06           ES 98         M00005449C:E10         ES 125         M0005642B:H08           ES 98         M00005449C:E10         E				<del></del>
ES 98				
ES 98	ES No.	Clone Name	ES No.	Clone Name
ES 98	ES 98	M00005309D:E05	ES 125	M00056371B:F12
ES 98	ES 98	M00005312A:D10	ES 125	M00056372B:C10
ES 98	ES 98	M00005313C:B02	ES 125	M00056374B:H02
ES 98	ES 98	M00005359A:A06	ES 125	M00056382C:H02
ES 98	ES 98	M00005373D:H06	ES 125	M00056383A:C10
ES 98	ES 98	M00005385C:A10	ES 125	M00056410B:E04
ES 98	ES 98	M00005389C:C11	ES 125	M00056411C:E04
ES 98	ES 98	M00005395A:D09	ES 125	M00056414B:A05
ES 98	ES 98	M00005406C:A11	ES 125	M00056416C:B12
ES 98	ES 98	M00005407B:E12	ES 125	M00056420C:D07
ES 98	ES 98	M00005411C:C07	ES 125	M00056421A:F12
ES 98	ES 98	M00005413B:F03	ES 125	M00056424A:F12
ES 98	ES 98	M00005415C:F12	ES 125	M00056424B:H06
ES 98	ES 98	M00005420B:C01	ES 125	M00056424D:A10
ES 98	ES 98	M00005438B:A06	ES 125	M00056425A:H08
ES 98	ES 98	M00005445C:A02	ES 125	M00056425D:B03
ES 98	ES 98	M00005447C:D01	ES 125	M00056434A:C08
ES 98	ES 98	M00005449C:E10	ES 125	M00056434D:E07
ES 98	ES 98	M00005454B:C03	ES 126	M00056480B:C12
ES 98	ES 98	M00005459B:B01	ES 126	M00056480D:A10
ES 98	ES 98	M00005469B:A07	ES 126	M00056481A:F02
ES 98	ES 98	M00005481D:C06	ES 126	M00056483D:F06
ES 98	ES 98	M00005485B:B05	ES 126	M00056484B:G02
ES 98	ES 98	M00005485D:A09	ES 126	M00056485B:B12
ES 98	ES 98	M00005491B:H12	ES 126	M00056490D:E02
ES 98	ES 98	M00005491D:B03	ES 126	M00056491D:G08
ES 98	ES 98	M00005500B:E03	ES 126	M00056496B:A01
ES 98	ES 98	M00005501B:E05	ES 126	M00056496C:H09
ES 98	ES 98	M00005501D:G09	ES 126	M00056499C:F05
ES 98	ES 98	M00005513B:F09	ES 126	M00056501C:H07
ES 98	ES 98	M00005514C:A06	ES 126	M00056503B:G11
ES 98	ES 98	M00005515B:H04	ES 126	M00056503B:G11
ES 98 M00005520C:E12 ES 126 M00056506C:G12 ES 98 M00005530C:A07 ES 126 M00056507D:B10 ES 98 M00005545B:A08 ES 126 M00056508B:B10 ES 99 M00005546A:G02 ES 126 M00056511A:H12 ES 99 M00005548A:A02 ES 126 M00056512B:C06 ES 99 M00005563C:B12 ES 126 M00056512C:E09 ES 99 M00005565A:F05 ES 126 M00056512D:C12 ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056515C:C05	ES 98	M00005516D:H06	ES 126	M00056505B:H02
ES 98 M00005530C:A07 ES 126 M00056507D:B10 ES 98 M00005545B:A08 ES 126 M00056508B:B10 ES 99 M00005546A:G02 ES 126 M00056511A:H12 ES 99 M00005548A:A02 ES 126 M00056512B:C06 ES 99 M00005563C:B12 ES 126 M00056512C:E09 ES 99 M00005565A:F05 ES 126 M00056512D:C12 ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 98	M00005517B:F04	ES 126	M00056505D:D07
ES 98	ES 98	M00005520C:E12	ES 126	M00056506C:G12
ES 99 M00005546A:G02 ES 126 M00056511A:H12 ES 99 M00005548A:A02 ES 126 M00056512B:C06 ES 99 M00005563C:B12 ES 126 M00056512C:E09 ES 99 M00005565A:F05 ES 126 M00056512D:C12 ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 98	M00005530C:A07	ES 126	M00056507D:B10
ES 99 M00005548A:A02 ES 126 M00056512B:C06 ES 99 M00005563C:B12 ES 126 M00056512C:E09 ES 99 M00005565A:F05 ES 126 M00056512D:C12 ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 98	M00005545B:A08	ES 126	M00056508B:B10
ES 99 M00005563C:B12 ES 126 M00056512C:E09 ES 99 M00005565A:F05 ES 126 M00056512D:C12 ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 99	M00005546A:G02	ES 126	M00056511A:H12
ES 99 M00005565A:F05 ES 126 M00056512D:C12 ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 99	M00005548A:A02	ES 126	M00056512B:C06
ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 99	M00005563C:B12	ES 126	M00056512C:E09
ES 99 M00005568C:B09 ES 126 M00056514B:E08 ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05		M00005565A:F05	<del></del>	<del></del>
ES 99 M00007926A:A07 ES 126 M00056514C:G01 ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 99	M00005568C:B09	ES 126	M00056514B:E08
ES 99 M00007926D:A05 ES 126 M00056515C:C05	ES 99	M00007926A:A07		M00056514C:G01
	ES 99	M00007926D:A05		M00056515C:C05
	ES 99	M00007927C:C01	ES 126	M00056517B:G03

Table 8	T	<del> </del>	T
ES No.	Clone Name	ES No.	Clone Name
ES 99	M00007931A:A10	ES 126	M00056519C:H01
ES 99	M00007935D:A05	ES 126	M00056526C:E11
ES 99	M00007936D:B09	ES 126	M00056529D:F12
ES 99	M00007936D:B09	ES 126	M00056529D:H09
ES 99	M00007939B:A03	ES 126	M00056530A:D01
ES 99	M00007943A:C02	ES 126	M00056532B:G06
ES 99	M00007951C:A05	ES 126	M00056534A:D11
ES 99	M00007953B:A01	ES 126	M00056537B:H05
ES 99	M00007953D:H09	ES 126	M00056537C:A09
ES 99	M00007954C:B04	ES 126	M00056541B:A08
ES 99	M00007961B:F05	ES 126	M00056547A:C04
ES 99	M00007964B:G01	ES 126	M00056548A:C11
ES 99	M00007965A:G10	ES 126	M00056551A:F02
ES 99	M00007965B:C03	ES 126	M00056552A:A10
ES 99	M00007965B:C03	ES 126	M00056552D:B10
ES 99	M00007981D:B04	ES 126	M00056555A:F09
ES 99	M00007982A:F11	ES 126	M00056556C:G01
ES 99	M00007983B:D03	ES 126	M00056557C:D02
ES 99	M00007983D:H06	ES 126	M00056561C:D08
ES 99	M00007990D:D03	ES 126	M00056564C:E09
ES 99	M00007991D:G01	ES 126	M00056566C:H01
ES 99	M00007992D:G08	ES 127	M00056574B:A07
ES 99	M00007994A:C11	ES 127	M00056580B:F10
ES 99	M00007994D:A05	ES 127	M00056591C:E03
ES 99	M00007998C:F07	ES 127	M00056592A:F04
ES 99	M00005589C:F07	ES 127	M00056592C:C03
ES 99	M00005610D:B11	ES 127	M00056592D:D07
ES 99	M00005619B:A09	ES 127	M00056592D:D07
ES 99	M00005621A:H08	ES 127	M00056593B:E05
ES 99	M00005627B:B10	ES 127	M00056594C:C06
ES 99	M00005628B:C10	ES 127	M00056594C:C10
ES 99	M00005632A:H02	ES 127	M00056595A:A02
ES 99	M00005650C:A06	ES 127	M00056595A:C07
ES 99	M00005650C:D04	ES 127	M00056595B:F02
ES 99	M00005655B:F08	ES 127	M00056596A:E02
ES 99	M00005675A:G02	ES 127	M00056596C:E06
ES 99	M00005685D:D12	ES 127	M00056596C:H08
ES 99	M00005704C:D10	ES 127	M00056597A:F07
ES 99	M00005708B:B07	ES 127	M00056597D:C02
ES 100	M00042455C:D11	ES 127	M00056599D:D11
ES 100	M00054826A:B05	ES 127	M00056600D:H07
ES 100	M00055281A:E08	ES 127	M00056603C:D01
ES 100	M00005657D:A12	ES 127	M00056608C:E04
ES 100	M00005710A:D01	ES 127	M00056610B:H12
ES 100	M00005765D:F07	ES 127	M00056613A:A05

Table 8	<del></del>		T
ES No.	Clone Name	ES No.	Clone Name
ES 100	M00005766C:F10	ES 127	M00056616B:C08
ES 100	M00005769B:A03	ES 127	M00056616D:A10
ES 100	M00005782A:B10	ES 127	M00056617B:H06
ES 100	M00005800D:D11	ES 127	M00056618A:B02
ES 100	M00005802B:H10	ES 127	M00056618B:F06
ES 100	M00005810B:F07	ES 127	M00056618D:F11
ES 100	M00005810B:G10	ES 127	M00056620D:E12
ES 100	M00005813B:E10	ES 127	M00056622D:C03
ES 100	M00005818D:B09	ES 127	M00056624D:H05
ES 100	M00005819C:B11	ES 127	M00056628C:F01
ES 100	M00005826D:G10	ES 127	M00056631B:G05
ES 100	M00005830C:D10	ES 127	M00056631D:C08
ES 100	M00006576A:B04	ES 127	M00056631D:D03
ES 100	M00006583B:H03	ES 127	M00056633B:B07
ES 100	M00006587B:A08	ES 127	M00056635A:A11
ES 100	M00006588A:H06	ES 127	M00056635A:E09
ES 100	M00006590A:C10	ES 127	M00056638A:D08
ES 100	M00006610C:D08	ES 127	M00056638B:B01
ES 100	M00006630D:C11	ES 127	M00056639A:E02
ES 100	M00006638A:G02	ES 127	M00056643D:G06
ES 100	M00006641C:H03	ES 127	M00056645C:B11
ES 100	M00006641C:H03	ES 127	M00056645D:F06
ES 100 ES 100	M00006648B:A05 M00006649D:B11	ES 127	M00056646C:C02 M00056646D:G05
ES 100	M00006650A:A03	ES 127	M00056652D:F04
ES 100	M00006650D:D05	ES 128	M00056656C:H03
ES 100	M00006664A:B09	ES 128	M00056659C:G08
ES 100	M00006679D:C04	ES 128	M00056661B:A09
ES 100	M00006686B:B07	ES 128	M00056661D:E05
ES 100	M00006695D:H08	ES 128	M00056662B:F03
ES 100	M00006695D:H08	ES 128	M00056664B:G06
ES 100	M00006704C:G06	ES 128	M00056664C:B07
ES 100	M00006705B:A09	ES 128	M00056665B:A11
ES 100	M00006705C:G09	ES 128	M00056665C:E05
ES 100	M00006712C:F02	ES 128	M00056666A:C08
ES 100	M00006719A:E12	ES 128	M00056669B:G07
ES 100	M00006719A:H07	ES 128	M00056670A:A11
ES 100	M00006731B:B02	ES 128	M00056673D:E06
ES 100	M00006731B:C08	ES 128	M00056674B:E05
ES 100	M00006731B:D03	ES 128	M00056674D:H04
ES 100	M00006731C:E01	ES 128	M00056682D:F10
ES 100	M00006734C:A08	ES 128	M00056683C:B09
ES 100	M00006737D:A11	ES 128	M00056684D:A05
ES 100	M00006740B:G01	ES 128	M00056684D:F11
ES 100	M00006745C:A02	ES 128	M00056688B:F05

Table 8	T	T	T
ES No.	Clone Name	ES No.	Clone Name
ES 100	M00006745D:D02	ES 128	M00056690C:F09
ES 100	M00006746C:B06	ES 128	M00056693C:C08
ES 101	M00006746C:B06	ES 128	M00056695A:H09
ES 101	M00006755D:A04	ES 128	M00056697C:E03
ES 101	M00006756B:F08	ES 128	M00056698C:E12
ES 101	M00006761C:C05	ES 128	M00056701B:E08
ES 101	M00006761C:D09	ES 128	M00056703A:D06
ES 101	M00006783B:F07	ES 128	M00056705D:E07
ES 101	M00006789D:A11	ES 128	M00056707B:E02
ES 101	M00006795D:A03	ES 128	M00056707D:D05
ES 101	M00006795D:D07	ES 128	M00056708C:C06
ES 101	M00006803A:C07	ES 128	M00056708D:D11
ES 101	M00006806B:C09	ES 128	M00056709A:A05
ES 101	M00006807A:G12	ES 128	M00056710A:C01
ES 101	M00006810A:D11	ES 128	M00056710B:F05
ES 101	M00006811D:D12	ES 128	M00056710B:H09
ES 101	M00006819B:B05	ES 128	M00056710D:F07
ES 101	M00006821D:B01	ES 128	M00056711A:C01
ES 101	M00006822C:A09	ES 128	M00056711A:F05
ES 101	M00006822D:E09	ES 128	M00056711A:F05
ES 101	M00006831D:H05	ES 128	M00056711D:A05
ES 101	M00006846A:D03	ES 128	M00056712C:A07
ES 101	M00006852A:C07	ES 128	M00056712C:B06
ES 101	M00006859B:D04	ES 128	M00056713D:G08
ES 101	M00006861B:A08	ES 128	M00056714C:H06
ES 101	M00006867C:F12	ES 128	M00056715A:D10
ES 101	M00006871D:F01	ES 128	M00056715A:E04
ES 101	M00006873C:E12	ES 129	M00056715A:G01
ES 101	M00006873D:B01	ES 129	M00056715B:C01
ES 101	M00006885C:G11	ES 129	M00056715D:C04
ES 101	M00006888A:G05	ES 129	M00056715D:E08
ES 101	M00006892B:F09	ES 129	M00056717B:C04
ES 101	M00006894D:A03	ES 129	M00056718C:B01
ES 101	M00006917B:A05	ES 129	M00056718C:G02
ES 101	M00006917C:A04	ES 129	M00056719A:D06
ES 101	M00006917D:D08	ES 129	M00056719A:F12
ES 101	M00006921A:H08 M00006923B:H05	ES 129	M00056719B:A09
ES 101	M00006923B:H03	ES 129 ES 129	M00056721A:F07 M00056722A:G01
ES 101	M00006928A:A04 M00006928B:D01	ES 129	M00056723B:D10
ES 101	M00006928B:D01	ES 129	M00036723B:D10
ES 101	M00006928B.D01 M00006937D:F03	ES 129	M00056724D:E11
ES 101	M00006957D.F03	ES 129	M00056726C:G05
ES 101	M00006967B:B05	ES 129	M00056728A:H05
ES 101	M00006976B:D05	ES 129	M00056728B:D05
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Table 8	Υ		T
ES No.	Clone Name	ES No.	Clone Name
ES 101	M00006979D:B10	ES 129	M00056729B:D04
ES 101	M00000977D:D10	ES 129	M00056729C:H12
ES 101	M00006989C:E04	ES 129	M00056733C:D09
ES 101	M00006990D:E02	ES 129	M00056735D:B08
ES 102	M00000996C:F10	ES 129	M00056737B:G07
ES 102	M00006997A:A03	ES 129	M00056739A:D11
ES 102	M00000997A:A03	ES 129	M00056739B:D08
ES 102	M00007003B:H05	ES 129	M00056740C:B05
ES 102	M00007028C:C04	ES 129	M00056741B:C06
ES 102	M00007028C:C04	ES 129	M00056746D:A02
ES 102	M00007028C:C04	ES 129	M00056746D:D06
ES 102	M00007032E.109	ES 129	M00056747A:D05
ES 102	M00007034B:B00 M00007035C:E06	ES 129	M00056752A:E01
ES 102	M00007055C:E00	ES 129	M00056753D:A10
ES 102	M00007084B:G04	ES 129	M00056754A:A04
ES 102	M00007092D:F03	ES 129	M00056754B:D09
ES 102	M00007096A:E02	ES 129	M00056754B:H04
ES 102	M00007096C:E01	ES 129	M00056754D:A05
ES 102	M00007096D:H02	ES 129	M00056756B:A05
ES 102	M00007097A:B04	ES 129	M00056756D:B08
ES 102	M00007097D:D07	ES 129	M00056757B:F03
ES 102	M00007098A:C05	ES 129	M00056758B:C05
ES 102	M00007105D:C12	ES 129	M00056759A:F11
ES 102	M00007108A:D01	ES 129	M00056759B:G03
ES 102	M00007110C:F03	ES 129	M00056760D:A04
ES 102	M00007112A:A12	ES 129	M00056761A:F05
ES 102	M00007117D:H03	ES 129	M00056762C:E05
ES 102	M00007121C:G08	ES 129	M00056763C:D05
ES 102	M00007128B:G06	ES 129	M00056764A:E08
ES 102	M00007129A:F08	ES 129	M00056765A:A10
ES 102	M00007131C:A01	ES 130	M00056765C:E12
ES 102	M00007135D:B11	ES 130	M00056765D:D10
ES 102	M00007135D:B11	ES 130	M00056766B:A10
ES 102	M00007136C:C05	ES 130	M00056771C:F12
ES 102	M00007146D:F11	ES 130	M00056771D:C12
ES 102	M00007151A:B11	ES 130	M00056772D:A04
ES 102	M00007156A:E06	ES 130	M00056772D:A04
ES 102	M00007156D:F08	ES 130	M00056772D:E08
ES 102	M00007166A:E06	ES 130	M00056773A:H11
ES 102	M00007172B:C03	ES 130	M00056774B:A02
ES 102	M00007174C:D06	ES 130	M00056775D:A07
ES 102	M00007177A:E11	ES 130	M00056775D:C01
ES 102	M00007178C:D03	ES 130	M00056775D:C08
ES 102	M00007192A:E06	ES 130	M00056776D:A06
ES 102	M00007194A:E06	ES 130	M00056776D:D09

ES No. Clone Name	TR. 1.1. 0	т	<del>                                     </del>	<del></del>
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ES 103 M00021649A:E12 ES 130 M00056811A:C04 ES 103 M00021654A:A04 ES 130 M00056812C:E08 ES 103 M00021654A:A04 ES 130 M00056815A:B01 ES 103 M00021655C:H02 ES 130 M00056816B:A10 ES 103 M00021670D:G05 ES 130 M00056817C:C03 ES 103 M00021681D:C02 ES 130 M00056821D:C09 ES 103 M00021681D:C02 ES 130 M00056821D:C09 ES 103 M00022189A:B03 ES 130 M00056823A:B05 ES 103 M00022216D:D06 ES 130 M00056823A:B05 ES 103 M0002221A:D06 ES 131 M00056824D:C10 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022251C:A09 ES 131 M00056840D:H09 ES 103 M00022253A:E03 ES 131 M00056841D:G09	ES 103	M00021628B:B11	ES 130	M00056805D:B09
ES 103 M00021654A:A04 ES 130 M00056812C:E08 ES 103 M00021655A:A04 ES 130 M00056815A:B01 ES 103 M00021655C:H02 ES 130 M00056816B:A10 ES 103 M00021670D:G05 ES 130 M00056817C:C03 ES 103 M00021681D:C02 ES 130 M00056821D:C09 ES 103 M00021681D:C02 ES 130 M00056821D:C09 ES 103 M00021681D:C02 ES 130 M00056822C:G11 ES 103 M00022189A:B03 ES 130 M00056823A:B05 ES 103 M00022216D:D06 ES 130 M00056823C:A07 ES 103 M0002221A:D06 ES 131 M00056824B:C10 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00021628B:D07	ES 130	M00056808B:B12
ES 103 M00021654A:A04 ES 130 M00056815A:B01 ES 103 M00021655C:H02 ES 130 M00056816B:A10 ES 103 M00021670D:G05 ES 130 M00056817C:C03 ES 103 M00021681D:C02 ES 130 M0005682D:C09 ES 103 M00021681D:C02 ES 130 M0005682C:G11 ES 103 M00022189A:B03 ES 130 M00056823A:B05 ES 103 M00022216D:D06 ES 130 M00056823C:A07 ES 103 M00022221A:D06 ES 131 M00056824B:C10 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022251C:A09 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00021649A:E12	ES 130	M00056811A:C04
ES 103 M00021655C:H02 ES 130 M00056816B:A10 ES 103 M00021670D:G05 ES 130 M00056817C:C03 ES 103 M00021681D:C02 ES 130 M00056821D:C09 ES 103 M00021681D:C02 ES 130 M00056821D:C09 ES 103 M00022189A:B03 ES 130 M00056823A:B05 ES 103 M00022216D:D06 ES 130 M00056823C:A07 ES 103 M0002221A:D06 ES 131 M00056824B:C10 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00021654A:A04	ES 130	M00056812C:E08
ES 103 M00021670D:G05 ES 130 M00056817C:C03 ES 103 M00021681D:C02 ES 130 M00056821D:C09 ES 103 M00021681D:C02 ES 130 M00056822C:G11 ES 103 M00022189A:B03 ES 130 M00056823A:B05 ES 103 M00022216D:D06 ES 130 M00056823C:A07 ES 103 M00022221A:D06 ES 131 M00056824D:E01 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022236A:H08 ES 131 M00056839A:G01 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00021654A:A04	ES 130	M00056815A:B01
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ES 103 M00021681D:C02 ES 130 M00056822C:G11 ES 103 M00022189A:B03 ES 130 M00056823A:B05 ES 103 M00022216D:D06 ES 130 M00056823C:A07 ES 103 M00022221A:D06 ES 131 M00056824B:C10 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022246A:H08 ES 131 M00056839A:G01 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00021670D:G05	ES 130	M00056817C:C03
ES 103 M00022189A:B03 ES 130 M00056823A:B05 ES 103 M00022216D:D06 ES 130 M00056823C:A07 ES 103 M00022221A:D06 ES 131 M00056824B:C10 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022236A:H08 ES 131 M00056839A:G01 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00021681D:C02	ES 130	M00056821D:C09
ES 103 M00022216D:D06 ES 130 M00056823C:A07 ES 103 M00022221A:D06 ES 131 M00056824B:C10 ES 103 M00022221D:D06 ES 131 M00056824D:E01 ES 103 M00022231D:E12 ES 131 M00056826A:B12 ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056830C:G02 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022236A:H08 ES 131 M00056839A:G02 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00021681D:C02	ES 130	M00056822C:G11
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ES 103 M00022234B:D05 ES 131 M00056830C:G02 ES 103 M00022235C:C11 ES 131 M00056833C:C01 ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022246A:H08 ES 131 M00056839A:G02 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00022221D:D06	ES 131	M00056824D:E01
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ES 103 M00022236A:A02 ES 131 M00056839A:G01 ES 103 M00022246A:H08 ES 131 M00056839A:G02 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00022234B:D05	ES 131	M00056830C:G02
ES 103 M00022246A:H08 ES 131 M00056839A:G02 ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00022235C:C11	ES 131	M00056833C:C01
ES 103 M00022251C:A09 ES 131 M00056839C:F01 ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00022236A:A02	ES 131	M00056839A:G01
ES 103 M00022253A:E03 ES 131 M00056840D:H09 ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00022246A:H08	ES 131	M00056839A:G02
ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	M00022251C:A09	ES 131	M00056839C:F01
ES 103 M00022259C:B07 ES 131 M00056841D:G09	ES 103	<del> </del>	+	<del></del>
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jes 105    M00022273B:A09	ES 103	M00022273B:A09	ES 131	M00056842B:F12
ES 103 M00022279B:H04 ES 131 M00056842B:F12	ES 103	M00022279B:H04	ES 131	M00056842B:F12

Table 8	1		T
ES No.	Clone Name	ES No.	Clone Name
ES 103	M00022280A:G11	ES 131	M00056843B:H09
ES 103	M00022280A:G11 M00022370A:G02	ES 131	M00056844A:E07
ES 103	M00022370A:G02	ES 131	M00056844C:A10
ES 103	M00022411D.G12 M00022415C:B06	ES 131	M00056848B:C07
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ES 103	M00022420B:H03	ES 131	M00056850B:E11
ES 103 ES 103	M00022430B:D10	ES 131	M00056850B:E11 M00056857B:C09
	M00022430C:D04 M00022440D:D01	ES 131	M00036857B:C09
ES 103	<del></del>	ES 131	<del></del>
ES 103	M00022444A:C11		M00056858B:A12
ES 103	M00022452A:B07	ES 131	M00056859A:D12
ES 104	M00022453B:H04	ES 131	M00056860A:F12
ES 104	M00022453B:H04	ES 131	M00056863C:E03
ES 104	M00022457C:G05	ES 131	M00056864B:H09
ES 104	M00022465D:F05	ES 131	M00056866B:E05
ES 104	M00022468A:E12	ES 131	M00056868D:E09
ES 104	M00022468C:E10	ES 131	M00056870A:E10
ES 104	M00022470B:G01	ES 131	M00056872A:A06
ES 104	M00022470B:G01	ES 131	M00056873C:E06
ES 104	M00022472D:E11	ES 131	M00056874B:H06
ES 104	M00022473D:B06	ES 131	M00056874C:D05
ES 104	M00022496D:F04	ES 131	M00056874D:G01
ES 104	M00022508A:C02	ES 131	M00056879A:E05
ES 104	M00022509A:B06	ES 131	M00056879B:H11
ES 104	M00022516B:E09	ES 131	M00056879D:A02
ES 104	M00022517B:E03	ES 131	M00056880D:B04
ES 104	M00022528A:H12	ES 131	M00056883D:A07
ES 104	M00022533C:E06	ES 131	M00056884B:C06
ES 104	M00022537A:C11	ES 131	M00056885C:C06
ES 104	M00022550C:B04	ES 131	M00056886A:C11
ES 104	M00022559D:D09	ES 131	M00056887B:F08
ES 104	M00022561A:A06	ES 131	M00056892C:A01
ES 104	M00022565A:A05	ES 131	M00056893B:H06
ES 104	M00022565A:A05	ES 131	M00056894D:G06
ES 104	M00022569A:A07	ES 131	M00056895B:A07
ES 104	M00022571C:D11	ES 131	M00056896A:F05
ES 104	M00021854C:E07	ES 131	M00056896A:F10
ES 104	M00021864A:E07	ES 132	M00056898D:D04
ES 104	M00021869D:D01	ES 132	M00056901A:A06
ES 104	M00021886D:F06	ES 132	M00056902A:H12
ES 104	M00021911A:H03	ES 132	M00056909B:E11
ES 104	M00021915B:E10	ES 132	M00056909C:D09
ES 104	M00021925C:H10	ES 132	M00056911B:F02
ES 104	M00021947B:C06	ES 132	M00056913B:G10
ES 104	M00022010B:H01	ES 132	M00056914D:B09
ES 104	M00022013D:H05	ES 132	M00056916C:B02

Table 8			<u> </u>
ES No.	Clone Name	ES No.	Clone Name
ES 104	M00022015D:F11	ES 132	M00056916C:F04
ES 104	M00022015D:F11	ES 132	M00056921A:C07
ES 104	M00022025C:D02	ES 132	M00056923C:E09
ES 104	M00022049C:B07	ES 132	M00056924D:B06
ES 104	M00022050C:D04	ES 132	M00056925D:C07
ES 104	M00022052D:A08	ES 132	M00056939A:F08
ES 104	M00022058D:A01	ES 132	M00056939D:B02
ES 104	M00022060B:F09	ES 132	M00056941D:E02
ES 104	M00022106B:D04	ES 132	M00056945A:B11
ES 104	M00022123A:D05	ES 132	M00056947D:F09
ES 104	M00022129A:E12	ES 132	M00056949C:F06
ES 104	M00022132A:D10	ES 132	M00056951B:F09
ES 104	M00022132C:F04	ES 132	M00056952C:A06
ES 105	M00022137B:G04	ES 132	M00056952D:H04
ES 105	M00022143A:C10	ES 132	M00056953B:A06
ES 105	M00022143A:D01	ES 132	M00056955B:G09
ES 105	M00022143A:D01	ES 132	M00056956B:F01
ES 105	M00022148A:A06	ES 132	M00056960A:C05
ES 105	M00022149C:C01	ES 132	M00056961A:B08
ES 105	M00022149D:C06	ES 132	M00056961C:G12
ES 105	M00022151A:D11	ES 132	M00056964B:A02
ES 105	M00022151A:G05	ES 132	M00056966D:A11
ES 105	M00022163A:C08	ES 132	M00056967A:D02
ES 105	M00022598B:E12	ES 132	M00056967A:E07
ES 105	M00022598C:D05	ES 132	M00056969B:C08
ES 105	M00022617B:C02	ES 132	M00056969D:B01
ES 105	M00022624C:C02	ES 132	M00056972A:F05
ES 105	M00022641A:C10	ES 132	M00056973D:B08
ES 105	M00022641A:E06	ES 132	M00056974C:F04
ES 105	M00022641B:F02	ES 132	M00056976C:F10
ES 105 ES 105	M00022645D:A05 M00022645D:C07	ES 132 ES 132	M00056977A:G03 M00056985B:C05
ES 105	M00022651D:B04	ES 132	M00056986A:F11
ES 105	M00022651D:B04	ES 132	M00056986D:G01
ES 105	M00022655A:D10	ES 132	M00056990C:B09
ES 105	M00022656D:E11	ES 132	M00056990D:C11
ES 105	M00022650D.E11	ES 132	M00056993A:B06
ES 105	M00022667A:C05	ES 132	M00056993D:D03
ES 105	M00022667D:E11	ES 132	M00056994B:F07
ES 105	M00022681D:E06	ES 132	M00056994C:C03
ES 105	M00022697A:D12	ES 133	M00056996D:A12
ES 105	M00022702B:B04	ES 133	M00056997C:H09
ES 105	M00022702B:B04	ES 133	M00056998A:E08
ES 105	M00022716C:C06	ES 133	M00057002D:B05
ES 105	M00022719A:F12	ES 133	M00057002D:B06
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Table 8	Γ		Γ
ES No.	Clone Name	ES No.	Clone Name
ES 105	M00022720B:A11	ES 133	M00057003B:B09
ES 105	M00022720B:711	ES 133	M00057005B:C01
ES 105	M00022724C:D04	ES 133	M00057005C:D03
ES 105	M00022724C:D04	ES 133	M00057007C:B12
ES 105	M00022741B:B11	ES 133	M00057008C:E09
ES 105	M00022741B:B11	ES 133	M00057000C:E03
ES 105	M00022749C:C07	ES 133	M00057013B:D01
ES 105	M00022750A:A07	ES 133	M00057015A:C12
ES 105	M00022791B:F11	ES 133	M00057019C:H02
ES 105	M00022813B:A08	ES 133	M00057023A:H09
ES 105	M00022820D:C06	ES 133	M00057024A:E02
ES 105	M00022823A:D03	ES 133	M00057024A:G05
ES 105	M00022828A:C06	ES 133	M00057024D:H08
ES 105	M00022829A:H06	ES 133	M00057025C:A08
ES 105	M00022829C:H10	ES 133	M00057027C:G06
ES 105	M00022831B:H07	ES 133	M00057028D:D09
ES 106	M00022831C:A09	ES 133	M00057029A:C12
ES 106	M00022831D:C04	ES 133	M00057029D:A06
ES 106	M00022834C:G01	ES 133	M00057033A:F09
ES 106	M00022836A:G03	ES 133	M00057035B:C09
ES 106	M00022853C:C11	ES 133	M00057041D:B11
ES 106	M00022861D:B10	ES 133	M00057044C:F06
ES 106	M00022872A:B05	ES 133	M00057047B:C02
ES 106	M00022876B:B05	ES 133	M00057049A:G06
ES 106	M00022876D:D08	ES 133	M00057049C:H05
ES 106	M00022880C:G09	ES 133	M00057052D:B11
ES 106	M00022892C:G07	ES 133	M00057052D:G09
ES 106	M00022895B:B11	ES 133	M00057055B:G08
ES 106	M00022897D:H03	ES 133	M00057055B:G08
ES 106	M00022898C:F04	ES 133	M00057058C:F09
ES 106	M00022899A:C09	ES 133	M00057059D:F06
ES 106	M00022901D:E11	ES 133	M00057059D:H09
ES 106	M00022901D:E11	ES 133	M00057060B:A12
ES 106	M00022902C:H10	ES 133	M00057061C:D04
ES 106	M00022908B:H03	ES 133	M00057063A:C08
ES 106	M00022911B:G01	ES 133	M00057065C:D04
ES 106	M00022928A:F03	ES 133	M00057066A:A04
ES 106	M00022934D:B03	ES 133	M00057070D:B08
ES 106	M00022956B:B09	ES 133	M00057072B:E02
ES 106	M00022961A:B11	ES 133	M00057073D:A05
ES 106	M00022973A:G07	ES 133	M00057074D:C09
ES 106	M00022973C:G08	ES 133	M00057074D:C09
ES 106	M00022974D:D10	ES 134	M00055909B:G01
ES 106	M00022995C:E02	ES 134	M00055909C:E08
ES 106	M00022997A:C08	ES 134	M00055911B:E06

Table 8	T		<del></del>
ES No.	Clone Name	ES No.	Clone Name
ES 106	M00022998B:C08	ES 134	M00055912C:E10
ES 106	M00022998B:C08	ES 134	M00055912D:C05
ES 106	M00023002D:G10	ES 134	M00055913B:D05
ES 106	M00023015A:D10	ES 134	M00055919A:A06
ES 106	M00023013C:D02	ES 134	M00055921A:E03
ES 106	M00023020D:G03	ES 134	M00055921B:B11
ES 106	M00023029A:E06	ES 134	M00055922A:C02
ES 106	M00023029A:E00	ES 134	M00055924A:H11
ES 106	M00023331D:A11	ES 134	M00055930A:B08
ES 106	M00023347D:C12	ES 134	M00055931A:A03
ES 106	M00023377D:C09	ES 134	M00055931A:C01
ES 106	M00023393D:E12	ES 134	M00055931B:E01
ES 106	M00023393D.E12	ES 134	M00055936B:E07
ES 106	M00023409A:G08	ES 134	M00055937B:C02
ES 106	M00023409A:G08	ES 134	M00035937B:C02
ES 106	M00023414B:F03	ES 134	M00055941B:B12
ES 106	M00023428C:D03	ES 134	M00055945A:H11
ES 106	M00023428D.F11	ES 134	M00055945B:E10
ES 106	M00023430B:D10	ES 134	M00055946D:G07
ES 100	M00023518C:A04	ES 134	M00055951C:C02
ES 107	M00026804D:D03	ES 134	M00055956C:E02
ES 107	M00026804B:B04	ES 134	M00055958D:F02
ES 107	M00026848C:G11	ES 134	M00055959D:A12
ES 107	M00026854A:E07	ES 134	M00055966C:A03
ES 107	M00026856C:C11	ES 134	M00055966C:D06
ES 107	M00026860D:E01	ES 134	M00055971C:E07
ES 107	M00026861D:A09	ES 134	M00055973A:D04
ES 107	M00026865D:G11	ES 134	M00055976B:F01
ES 107	M00026866A:H08	ES 134	M00055979B:B09
ES 107	M00026873B:E11	ES 134	M00055980A:A10
ES 107	M00026873D:B08	ES 134	M00055981D:A07
ES 107	M00026879A:B02	ES 134	M00055984C:C02
ES 107	M00026879C:D10	ES 134	M00055985D:D01
ES 107	M00026890C:D02	ES 134	M00055990C:B05
ES 107	M00026893C:A01	ES 134	M00055992C:E11
ES 107	M00026896D:E10	ES 134	M00056139D:E04
ES 107	M00026899C:G11	ES 134	M00056139D:G01
ES 107	M00026899C:G11	ES 134	M00056140B:H07
ES 107	M00026900B:C02	ES 134	M00056140D:E07
ES 107	M00026902A:G04	ES 134	M00056141A:D05
ES 107	M00026906B:C10	ES 134	M00056141D:B09
ES 107	M00026909A:G03	ES 134	M00056143A:E09
ES 107	M00026917D:H03	ES 134	M00056144B:C09
ES 107	M00026926D:C05	ES 134	M00056145C:B04
ES 107	M00026934D:E09	ES 134	M00056149C:B01

Table 8	T	<del></del>	1
ES No.	Clone Name	ES No.	Clone Name
ES 107	M00026936D:C12	ES 135	M00056150B:C12
ES 107	M00026937C:B08	ES 135	M00056153C:D01
ES 107	M00026938A:F04	ES 135	M00056156D:A12
ES 107	M00026938A:F04	ES 135	M00056160D:A08
ES 107	M00026949B:H10	ES 135	M00056161D:G04
ES 107	M00026950A:F12	ES 135	M00056162B:F08
ES 107	M00026950D:H01	ES 135	M00056162B:F08
ES 107	M00026951A:G06	ES 135	M00056162D:D06
ES 107	M00026951A:G11	ES 135	M00056162D:E09
ES 107	M00026951A:G11	ES 135	M00056167D:B08
ES 107	M00026951C:D03	ES 135	M00056169A:F06
ES 107	M00026975C:B03	ES 135	M00056171C:H11
ES 107	M00026977A:E09	ES 135	M00056171C:H12
ES 107	M00026984A:D10	ES 135	M00056180B:H09
ES 107	M00026985C:B05	ES 135	M00056184B:D08
ES 107	M00026986B:H10	ES 135	M00056184C:H03
ES 107	M00026993B:H06	ES 135	M00056184D:F01
ES 107	M00026994C:A07	ES 135	M00056185D:A03
ES 107	M00026996D:A06	ES 135	M00056185D:D06
ES 107	M00027000C:F05	ES 135	M00056186C:F02
ES 107	M00027006B:H01	ES 135	M00056190D:G02
ES 107 ES 108	M00027013D:E10 M00027014C:G04	ES 135 ES 135	M00056192D:E04
ES 108	M00027014C:G04 M00027014D:G04	ES 135	M00056192D:H02 M00056195B:C08
ES 108	M00027014D:G04	ES 135	M00056198A:D07
ES 108	M00027010D:H11	ES 135	M00056199D:A09
ES 108	M00027028D:C07	ES 135	M00056201C:H08
ES 108	M00027030C:C08	ES 135	M00056203A:H10
ES 108	M00027034B:D09	ES 135	M00056204B:A04
ES 108	M00027034C:D11	ES 135	M00056205B:D01
ES 108	M00027035D:H09	ES 135	M00056206A:E06
ES 108	M00027039A:F06	ES 136	M00055997C:G11
ES 108	M00027039B:E09	ES 136	M00055999C:G10
ES 108	M00027042C:G11	ES 136	M00055999D:G06
ES 108	M00027046B:E05	ES 136	M00056000A:F12
ES 108	M00027051A:A07	ES 136	M00056000C:D09
ES 108	M00027054B:B03	ES 136	M00056001A:B06
ES 108	M00027076D:F07	ES 136	M00056001A:B07
ES 108	M00027084C:H10	ES 136	M00056001C:E09
ES 108	M00027088D:H06	ES 136	M00056003A:E06
ES 108	M00027090A:E08	ES 136	M00056005B:E05
ES 108	M00027093C:B08	ES 136	M00056005D:C04
ES 108	M00027096A:G07	ES 136	M00056007A:A11
ES 108	M00027097C:G11	ES 136	M00056007C:F06
ES 108	M00027111A:H04	ES 136	M00056016D:D06

Table 8	T		T
ES No.	Clone Name	ES No.	Clone Name
ES 108	M00027134A:G02	ES 136	M00056018B:G05
ES 108	M00027139D:C06	ES 136	M00056020A:D10
ES 108	M00027140A:C11	ES 136	M00056020D:D07
ES 108	M00027163A:D11	ES 136	M00056028C:F03
ES 108	M00027165C:F11	ES 136	M00056036D:B06
ES 108	M00027168C:H10	ES 136	M00056037C:B02
ES 108	M00027171D:B07	ES 136	M00056038D:F04
ES 108	M00027172A:C03	ES 136	M00056041A:C04
ES 108	M00027173D:D08	ES 136	M00056042A:A01
ES 108	M00027183B:B01	ES 136	M00056045D:H01
ES 108	M00027193C:C05	ES 136	M00056050C:A03
ES 108	M00027194D:A05	ES 136	M00056053A:A09
ES 108	M00027194D:A03	ES 136	M00056053A:A09
ES 108	M00027197A:G07	ES 136	M00056055A:A07
ES 108	M00027197B:F07	ES 136	M00056055B:B01
ES 108	M00027207B:E09	ES 136	M00056055C:D03
ES 108	M00027207B:E03	ES 136	M00056058A:H04
ES 108	M00027217A:G03	ES 136	M00056060B:B10
ES 108	M00027220A:B12	ES 136	M00056061B:F06
ES 108	M00027222A:C09	ES 136	M00056066D:H07
ES 108	M00027229D.E00	ES 136	M00056067B:D08
ES 108	M00027524B:B11	ES 136	M00056074D:G10
ES 108	M00027527A:G04	ES 136	M00056077D:E06
ES 108	M00027527A:G04 M00027532C:C02	ES 136	M00056077D:E12
ES 108	M00027535D:E08	ES 136	M00056077D:E12
ES 109	M00027535D:E08	ES 136	M00056079B:D12
ES 109	M00027543C:B09	ES 136	M00056079B:F07
ES 109	M00027543D:G07	ES 136	M00056079C:C11
ES 109	M00027556D:G10	ES 136	M00056081D:B05
ES 109	M00027561C:C04	ES 136	M00056081D:B09
ES 109	M00027562B:C02	ES 136	M00056082C:F06
ES 109	M00027564A:D03	ES 136	M00056085D:H11
ES 109	M00027571C:C11	ES 136	M00056094A:H07
ES 109	M00027573A:F09	ES 136	M00056098A:H01
ES 109	M00027578B:F05	ES 136	M00056099B:G09
ES 109	M00027578C:E04	ES 136	M00056099B:H11
ES 109	M000275780C:E10	ES 136	M00056099B:H11
ES 109	M00027581B:E01	ES 136	M00056103A:D12
ES 109	M00027588A:C01	ES 136	M00056103C:H12
ES 109	M00027588C:A06	ES 136	M00056107B:E06
ES 109	M00027588C:A00	ES 136	M00056108D:B12
ES 109	M00027594B.C03	ES 136	M00056108D:B12
ES 109	M00027604A:G10	ES 136	M00056110C:D09
ES 109	M00027605C:E05	ES 136	M00056111D:H02
ES 109	M00027607A:H05	ES 136	M00056112A:H02
103	1M10002/00/A.TU3	E9 130	1M100030112/A.IIUZ

Table 8			
ES No.	Clone Name	ES No.	Clone Name
ES 109	M00027608C:H07	ES 136	M00056114C:C06
ES 109	M00027616C:G12	ES 136	M00056125B:D09
ES 109	M00027628C:A01	ES 136	M00056128C:B10
ES 109	M00027639B:E11	ES 136	M00056131B:C12
ES 109	M00027641B:A01	ES 136	M00056133D:D09
ES 109	M00027652B:G03	ES 136	M00056136A:B11

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Comment	invasive adenocarcinom a, moderately differentiated; focal perineural invasion is seen	Hyperplastic	polyp m appendix.		Perineural invasion; donut	anastomosis	negative. One		patient history	of metastatic	melanoma												
Dist Met Grade	×			W0				₩							OV.				M0			M	
Descrip Distant Met													_	_									
Distant Met & Loc		21112		negative				negative							.,	IICganve			negative			negative	,
Region al Lymph	7			N0				N0								ON			ī			NZ	-
Incidenc Region e al Lymphn Lymph	Ç	5		0/12				0/34							0.50	61/0			1/5			10/24	
Lymphno I de Met		positive		negative				negative							•	negative			positive			nosifive	_
Histopath Local Invasion Lymphno Incidenc Region Grade de Met e al Lymphn Lymph	extending into subserosal adipose tissue	Invasion	through muscularis	ргорпа,	Invasion of	propria into	serosa,	gurvlovni	Invasion	through the	muscularis	propria into	suserosal	adipose tissuc.	Ileocccal	Janenou:	Invasion of	propria into	percolonic fat	through wall	and into	surrounding admose tissue	- I dian
Histopath Grade		75		3				C5								3			C5				
Primary Tumor Grade		113		T3				T4	*							T.							112
Primary Tumor Size		4.0		0.6				9								9			0.5	3		ų	CC
Anatomical Loc	Ascending	colon	A contract	colon				Sigmoid	Signion							Cecum		E	Iransverse	10103		Splenic	tlexure
Path Report ID		21		71				140	1							144			147	Ì			149
Table 9 PatientID		15		63	35			121	121							125			92	071	w <del>r.</del>		130

## girth girth girth grad, grad, grad, grad, det gerth, grad, grad, grad, grad, grad, grad, grad, grad, sall, land sall, sall fladf sall, sal

Comment	Small separate tubular adenoma (0.4 cm)	Perineural invasion identified adjacent to metastatic adenocarcinom a.	Separate tubolovillous and tubular adenomas
Dist Met Grade	MO	W	M0
Descrip Distant Met		adenocarcin oma consistant with	
Distant Met & Loc	negative	positive (Liver)	negative
Region al Lymph	OZ	CZ	Z
Incidenc Region e al Lymphn Lymph	6/0	127	2/13
Lymphno   I de Met 	negative	pocifive	positive
Histopath Local Invasion Lymphno Incidenc Region  Grade de Met e al  Lymphn Lymph	Invasion through muscularis propria into non- peritonealized pericolic tussue, gross configuration is annular.	Invasion of muscularis propria into pericolonic adipose tissue, but not through scrosa. Arising from tubular adenoma.	Invasion through mucsularis propria into subserosa/peric olic adipose, no serosal involvement. Gross configuration annular.
Histopath Grade	8	8	5
Primary Tumor Grade	Ę	3	E E
Primary Tumor Size	C		3.8
Anatomical Loc			Lecum Hepatic flexure
Path Report ID		301	175
Table 9 PatientID	5	CCI	156

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Comment		Hyperplasiic polyps					Tubulorillous	adenoma with	hioh orade	dvsnlasia	nichide (n						_												:	Descending	colon polyps,	no HGD or	carcinonia	
Dist Met Grade	—I⊢	<u> </u>			,	XX	VIII.					-	MIO								,	XX						3	₽ E				9	MO
Descrip Distant Met	Talke															0.4 cm,	may	represent	lymph node	completely	Mesenteri replaced by	tumor											<u>-</u>	
Distant Met &	LOC					.,	neganve						negative							positive	(Mesenter	c deposit) tumor							negative					negative
Region al	Lympii					-	ž	. "					02									z							2					Z2
ncidenc e	Lympnn Lympu					9	8/-						01/0									0/15							0/12					2/10
Jymphno I de Met	-						positive	-					negative									negative							negative					positive
Histopath Local Invasion Lymphno Incidenc Region Grade e al		Invasion through muscularis	propria to	subserosal,	perirectoal	adipose, and	SCIOS	Invasion	through	muscularis	propria into	subscrosal	adipose tissuc	Invades through	muscularis	propria to	involve	pericolonic	adipose,	extends to	serosa.		Invades full	thickness of	muscularis	propria, but	mesenteric	adipose mec or	malignancy	Invasion into	perirectal	adipose tissuc.		
Histopath Grade			_				G2 to G3						G2									CZ							C5					G2
	Grade						T3						13									T3							12				_	T3
Primary Tumor Size							5.8						5.5									6							6.5					4
Anatomical Loc	. 1		•				Rectum					Ascending	colon								Transverse	colon							Cecum					Rectum
-	<u> </u>						247						283									285							287					297
Table 9 PatientID							228						264									3,66							368					278

#### ուրը ընտ կարել դրար դրար եր դրա չիրու դեր ըրուցիության հայի դրար դրար արեր գրուցիության ուրը արեր դրար դրար ար Table 9

Comment	Melanosis coli and diverticular disease.	I hyperplastic polyp identified		To see	polyps	at prior ileocolic surgical anastomosis.
Dist Met Grade	MO do a	M0	MX	M0	M0	M1
Descrip Distant Met (						Macrovesic ular and microvesic ular steatosis
Distant Met & Loc	negative	negative	negative	negative	negative	positive (Liver)
Region al Lymph	S	0N	°Z	0N	Z	- Z
Incidenc Region e al Lymphn Lymph	6170	9/0	0/4	0/4	1/5	9/1
Jymphno II de Met		negative	negative	negative	positive	positive
Histopath Local Invasion Lymphuo Incidenc Region Grade de Met e al Lymphn Lymph	Invasion through muscularis propria into percolic adipose	Extends into perirectal fat but does not reach serosa	Invasion through muscularis propria to involve pericolonic fat. Arising from villous adenoma.	Through colon wall into subserosal adipose tissue No serosal spread seen.	Invasion thru muscularis propria to pericolonie fat	Invasion through muscularis propria into subserosal adipose tissue, not serosa.
Histopath Grade		63	8	G2	G2	62
Primary Tumor		ET E	: :	2 [2	£	T
Primary Tumor Size		5.0	2 cm	5.9	2, 4,3	
Anatomical Loc	Ascending	colon Rectosigmo	Ascending	Cotton	Ascending	Ascending
_ t	2	314	8000	360	515	444
Table 9 PatientID		295	339	341	350	392

## 

Comment									-	ediagnosis of	conforectomy	nath to	metastatic	colon cancer.					Anatonnear location of	primary not	notated in	report.	Evidence of	chronic colitis.				Mo montion of	No incinion of	distant met in	report			
Dist Met Grade									9	$\top$							7	IW										Ξ					9	OM
Descrip Distant Met															adenocarcin	oma in	multiple	slides		•	moderately	differentiat	B	adenocarcin	oma,		with	primary						
Distant Met & Loc										negative							positive	(Liver)									positive	(Liver)						negative
Region al Lymph										0Z								9 2										Z			_			N2
Incidenc Region  e al I ymphn Lymph	Junpun 6									0/21								2/0			_				_			2/17						9/9
ymphno li de Met	<b>-</b>									negative								negative						_	_			positive						positive
Histopath Local Invasion Lymphno Incidenc Region Grade de Met e al Local Lymph		Cecum, invades	through	muscularis	propria to	involve	subserosal	adipose tissuc	but not scrosa.		Invasive	through	muscularis to	involve	periserosal fat,	abutting	ileocecal	junction.	Invasion	through	muscularis	ргорга	guivlovui	bericolic	adipose, scrosal	surface	nninvolved		nenetrates	muscularis	propria,	involves	pericolonic fat	
Histopath Grade										Q5								CZ						_				G5			_			C5
1.	Grade									T3								Ţ.																T3
Primary Tumor Size						_			_	0.9			_		•			0 7	2								į	/.5 cm max						3
Anatomical Loc										Cacum							Accepting	Ascending	LOIOI															Sigmoid
Path A	E								_	344	£								402	·									585					395
Table 9											393								413										505					517

# 

Comment	Omentum with fibrosis and fat necrosis. Small bowel with acute and chronic serositis, focal abscess and adhesions.			Appendix dilated and fibrotic, but not involved by tumor
Dist Met Grade	0 4 1 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	M0	M	MO
Descrip Distant Met			metastatic adenocarcin oma	
Distant Met & Loc		negative	positive (Liver)	negative
Region al Lymph		0Z	Z 	0 <u>V</u>
Incidenc Region e al Lymphn Lymph		8/0	6/12	
Jymphno I de Met		negative	Pocultivo	
Histopath Local Invasion Lymphno Incidenc Region  Grade de Met e al Lymphn Lymphn Lymphn Lymphn	Invasion through the muscularis propria involving pericolic fat Serosa free of tumor		Invasion through muscularis propria extensively through submucosal and extending to serosa.	Invasion through the bowel wall, mto suberosal adipose. Serosal surface free of tumor
Histopath Grade		83		3 3
Primary Tumor Grade		T3		E E
Primary Tumor Size		12		5.5
Anatomical Loc		Ascending	Ascending	colon
Path A		553		565
Table 9 PatientID		534		546

# 

Comment	tubular	adenoma and hyperplstic	polyps present,	moderately differentiated	adenoma with	mucinous	(% not stated)		invasive poorly	adenosquamou	s carcinoma		moderately	differentiated	invasive	adenocarcinom	8			poorly	differentiated	invasive	colonic	adenocarcinom	a	
Dist Met Grade								MX				MI	_						M			_			M	
Descrip Distant Met															_											
Distant Met & Loc								negative			nositive	(Liver)							positive (1 iver)					positive	(Liver)	
Region al Lymph		_		_								NZ						_	2	ON					NZ	
Incidenc Region e al Lymphn Lymph								0/22				2/17	5							71/0					13/25	
de Met								nogative	a magni			ocitive.	DOSHING							negative					positive	-
Histopath Local Invasion Lymphuo Incidenc Region  Grade e de Met e al  Lymphn Lymphn Lymphn		extending through bowel	wall into scrosal	Iai	_				through	muscularis	propria into	pencone son	CONCEI	through	muscularis	propria into	pericolic fat,	but not at	serosal surface		through the	muscularis	propria into	pericolic tat		
Histopath   Grade								(	3				3							C5						3
Primary Tumor Grade	,							;	T3				IJ							T3						13
Primary Tumor Size			_						14				3.5							9.5						5.8
Anatomical Loc	_								Cecum			Ascending	colon						Descending	colon					Ascending	colon
_ =	<b>a</b>								714				803	3						808						810
Table 9 PatientID									695				787	10/						786	00/					162

# The first part hard then then the first the term for the first benefits that the benefits the term to the first term term to the first ter

Table 9 Path A	Anatomical Loc	Primary Tumor Size	Primary Tumor Grade	Histopath Grade	Histopath Local Invasion Lymphno Incidenc Region Distant Grade dc Met e al Met & Met & Doc	Lymphno de Met	Incidenc Region D  e al N  Lymphn Lymph	Region al Lymph	Distant Met & Loc	Descrip Distant Met	Dist Met Grade	Comment
=								-				well- to
					into muscularis							moderately-
					ргорпа		,					differentiated
												adenocarcinom
												a; this patient
												has tumors of
												the ascending
					_							colon and the
												sigmoid colon
									positive			)
	Ascending		i	-		positive	3/21	0X	(Liver)		Ξ	
806	colon	2.0	12	5		hone						moderately
					through				positive			differentiated
					muscularis	_	7/1	Z	(Tiver)		M	adenocarcinom
000	C. C. C.	2.7	<u>T</u>	3	propria int	positive		1 1	(1)			

Table 10

	%Pts	% Pts	% Pts				
SEQ ID	>=2x	>=2_5x	>=5x				
NO	T/N	T/N	T/N	P15	P52	P121	P125
18	30.3	15.2	3.0	1.855	2.705	1.000	2.280
22	45.5	39.4	18.2	2.196	1.719	0.604	2.388
127	27.3	18.2	6.1	1.000	1.620	1.822	1.692
139	21.2	18.2	15.2	1000.000	0.001	2.345	1.000
148	27.3	18.2	6.1	1.000	1.620	1.822	1.692
155	45.5	12.1	3.0	1.870	3.104	1.361	2.388
246	42.4	9.1	0.0	2.211	2.347	1.000	1.493
272	48.5	27.3	12.1	1.735	3.110	1.379	2.277
273	21.2	18.2	18.2	1.000	1.000	0.330	1.349
279	24.2	12.1	0.0	1.614	2.348	1.498	1.916
298	21.2	18.2	18.2	1.000	1.000	0.330	1.349
329	21.2	9.1	6.1	1.000	1.000	2.211	1.182
353	45.5	12.1	3.0	1.870	3.104	1.361	2.388
354	48.5	30.3	3.0	1.000	1.592	2.248	2.315
381	27.3	18.2	6.1	1.000	1.620	1.822	1.692
389	21.2	9.1	6.1	1.000	1.000	2.211	1.182
405	21.2	9.1	3.0	1.000	2.366	1.546	1.562
406	21.2	9.1	3.0	1.000	2.366	1.546	1.562
412	36.4	18.2	0.0	2.584	1.332	1.952	1.641
421	51.5	24.2	3.0	2.481	2.253	2.234	1.431
465	21.2	18.2	15.2	1000.000	0.001	2.345	1.000
534	21.2	9.1	3.0	1.000	2.366	1.546	1.562
538	42.4	15.2	0.0	1.489	2.019	3.022	1.121
565	45.5	12.1	3.0	1.870	3.104	1.361	2.388
657	45.5	30.3	3.0	1.512	2.748	0.784	2.162
670	24.2	6.1	0.0	1.190	1.000	0.656	1.456
739	21.2	12.1	0.0	1.936	1.830	0.831	1.347
741	48.5	18.2	0.0	2.750	2.458	1.485	1.151
744	48.5	21.2	0.0	2.069	3.002	1.229	1.631
755	30.3	18.2	3.0	1.000	1.414	1.236	1.738
757	21.2	15.2	6.1	1.000	0.839	2.032	2.557
810	30.3	18.2	3.0	1.000	1.414	1.236	1.738
811	30.3	18.2	3.0	1.000	1.414	1.236	1.738
845	30.3	15.2	9.1	1.000	0.271	0.860	1.310
861	24.2	21.2	15.2	1000.000	1000.000	1.000	1.320
915	30.3	15.2	3.0	1.855	2.705	1.000	2.280
954	24.2	21.2	15.2	1000.000	1000.000	1.000	1.320
955	39.4	21.2	3.0	1.612	2.281	0.785	2.045
991	39.4	21.2	3.0	1.612	2.281	0.785	2.045
1035	24.2	6.1	0.0	1.190	1.000	0.656	1.456
1049	21.2	12.1	0.0	1.936	1.830	0.831	1.347
1050	39.4	21.2	3.0	1.612	2.281	0.785	2.045
1235	45.5	12.1	3.0	1.870	3.104	1.361	2.388
1292	21.2	3.0	0.0	1.558	2.014	2.250	1.643

Table 10

	%Pts	% Pts	% Pts		T			
SEQ ID	>=2x	>=2_5x	>=5x					
NO	T/N	T/N	T/N	P15		P52	P121	P125
1313	24.2	6.1	0.0	1.19	0	1.000	0.656	1.456
1331	27.3	9.1	3.0		:7	3.749	1.000	2.045
1334	48.5	30.3	3.0		0	1.592	2.248	2.315
1418	30.3	15.2	3.0		55	2.705	1.000	2.280
1419	45.5	12.1	3.0		70	3.104	1.361	2.388
1420	39.4	18.2	3.0		59	1.566	1.000	2.302
1477	39.4	18.2	3.0		59	1.566	1.000	2.302
1554	33.3	15.2	0.0	1.83	29	1.622	1.882	1.957
1579	42.4		0.0	2.2	11	2.347	1.000	1.493
1600	45.5		3.0	1.8	70	3.104	1.361	2.388
1639	48.5			1 1.7	35	3.110	1.379	2.277
1657	42.4				00	1.908	2.267	1.188
1679				0 1.8	29	1.622	1.882	1.957
1744		<b></b>			59	2.753	1.679	1.587
1847					25	4.493	1.642	2.743
1877					70	3.104	1.361	2.388
1880					00	4.075	1.754	
1889					370	3.104	1.361	
2009					355	2.705	1.000	
2009					285	2.400	0.767	
2025					211	2.347	1.000	
2071		<del></del>			190	1.000	0.656	
207			<u> </u>		190	1.000	0.656	
2103					505	0.793	0.809	
210					190	1.000		
213					000	0.296		
213					263	1.000		
$\frac{214}{218}$					945	2.010		
218				3.0 1.	000	1.414	1.23	
219					190	1.000		
220				3.0 1	.870	3.104		
220	<del></del>			3.0 2	.256	2.228		
222					.591	0.483		
222					.602	3.209		
224					.870	3.10	4 1.36	
225	·				.985	2.26		
220					.886	1.00		
220					.529	3.04		
22			5.4		.008	0.68		
22			5.2		.855	2.70	5 1.00	
23			5.2		.855	2.70		
23			9.1		2.211	2.34		
	<u> </u>		1.2		2.497	1.83		
			8.5		2.603	2.64	1.0	00 1.93

Table 10

	%Pts	% Pts	% Pts				
SEQ ID	>=2x	>=2_5x	>=5x				
NO	T/N	T/N	T/N	P15	P52	P121	P125
2314	48.5	27.3	12.1	1.735	3.110	1.379	2.277
2316	42.4	9.1	0.0	2.211	2.347	1.000	1.493
2327	39.4	24.2	3.0	2.006	1.692	1.778	1.662
2348	72.7	45.5	0.0	2.961	3,152	2.712	1.346

Table 11

SEQ ID				D1.41	P156	P228	P264	P266
NO	P128	P130	P133	P141		1.457	2.262	1.236
18	0.713	1.800	1.955	0.663	0.466 1.000	2.647	1.628	1.190
22	1.594	6.800	1.340	1.131		1.000	1.000	1.000
127	3.761	1.000	1.000	1.587	2.127	0.767	1.631	1.000
139	1000.000	1.000	1000.000	0.482	2.846	1.000	1.000	1.000
148	3.761	1.000	1.000	1.587	2.127	1.306	2.099	1.357
155	2.062	1.781	2.302	1.000	1.000	1.301	2.098	1.733
246	1.779	1.337	2.865	1.515	1.617		1.588	1.634
272	2.044	2.219	4.257	0.744	1.000		1.000	1.000
273	1000.000	1000.000	1.000	1.000	0.566		1.399	1.239
279	1.202	1.852	2.370	1.000	1.000		1.000	1.000
298	1000.000		1.000	1.000	0.566		0.001	1.445
329	3.234		1.000	8.480	2.077			1.357
353	2.062	1.781	2.302	1.000	1.000			2.059
354	1.664		2.307	2.728	1.000			1.000
381	3.761		1.000	1.587	2.127		<del></del>	1.445
389	3.234		1.000	8.480	2.077			1.64
405	1.531	1	1.854	2.044	1.363			1.64
406	1.531		1.854		1.363			2.019
412	1.831		2.326		1.773			2.25
421	2.209		3.114					1.00
465	1000.000		1000.000		2.840			1.64
534	1.531		1.854					2.48
538	1.559		1.740					1.35
565	2.062		2.302					2.26
657	1.524		2.846					0.77
670	1.182				<u> </u>		<u> </u>	1.78
739	0.84							
741	1.81		3.22					
744	2.51							
755								
757								
810								
811								
845								
861								
915								
954								
955							-	
991								
1035					4			
1049								
105								
123								
129						$\frac{1.2}{000}$		
131	3 1.1	82 1.63	6 1.4	18 1.29	1.0	1.0	<u> </u>	

Table 11

						1 000	1.667	1.000	1.374
	1331	1.427	1.669	1.837	1.265	1.000	1.239	1.469	2.059
	1334	1.664	1.987	2.307	2.728	0.466	1.457	2.262	1.236
	1418	0.713	1.800	1.955	0.663	1.000	1.306	2.099	1.357
	1419	2.062	1.781	2.302	1.000	1.000	1.234	1.186	1.730
	1420	1.518	1.997	2.298	2.273		1.234	1.186	1.730
	1477	1.518	1.997	2.298	2.273	1.000	1.000	2.042	1.970
	1554	2.959	1.821	2.234	1.181	1.617	1.301	2.098	1.733
	1579	1.779	1.337	2.865	1.515	1.000	1.306	2.099	1.357
	1600	2.062	1.781	2.302	1.000	1.000	1.127	1.588	1.634
	1639	2.044	2.219	4.257	0.744	2.974	1.798	1.899	2.065
	1657	2.160	1.416	1.000	3.531	1.827	1.000	2.042	1.970
	1679	2.959	1.821	2.234	1.181	1.367	1.605	2.145	2.098
	1744	1.479	1.669	2.442	1.352	1.816	1.352	3.390	2.541
	1847	1.839	2.548	2.954	2.234	1.000	1.306	2.099	1.357
	1877	2.062	1.781	2.302	1.000	2.391	1.675	2.572	3.031
	1880	2.762	2.081	4.111	2.306	1.000	1.306	2.099	1.357
	1889	2.062	1.781	2.302	1.000	0.466	1.457	2.262	1.236
	2009	0.713	1.800	1.955	0.663	1.718	1.197	1.965	2.023
	2023	1.871	1.869	2.588	1.834	1.617	1.301	2.098	1.733
	2029	1.779	1.337	2.865	1.515	1.000	1.000	1.127	0.774
	2071	1.182	1.636	1.418	1.298	1.000	1.000	1.127	0.774
	2077	1.182	1.636	1.418	1.607	0.252	1.602	2.866	1.292
	2103	2.297	0.855	1.659	1.298	1.000	1.000	1.127	0.774
	2109	1.182	1.636	1.418	2.403	0.647	0.605	0.469	0.528
	2138	2.074	1.438	1.552	2.548	0.073	1.201	1.722	1.181
	2143	2.828	2.795	2.732	1.688	1.230	1.241	1.237	1.852
	2183	1.714	3.061	4.635 2.234	3.723	1.000	1.285	1.771	2.246
	2185	1.000	0.754	1.418	1.298	1.000	1.000	1.127	0.774
L	2190	1.182	1.636	2.302	1.000	1.000	1.306	2.099	1.357
L	2200	2.062	1.781	2.302	2.235	2.121	1.388	3.468	2.115
L	2206	2.229	2.131	1.629	1.586	0.155	1.408	2.830	1.636
	2220	2.650	0.815	2.510	0.628	1.763	1.000	1.000	1.687
	2224	1.385	2.044	2.302	1.000	1.000	1.306	2.099	1.357
	2249	2.062	1.781	1.567	2.350	1.729	2.071	1.439	1.540
	2255	1.454	2.931	1.690	1.678	0.724	2.656	2.035	3.526
	2265	2.843	1.937	3.729	2.105	2.224	2.547	2.605	4.402
Į	2267	2.490	2.374	1.404	4.761	3.241	2.253	1.384	1.912
1	2281	3.412	1.800	1.955	0.663	0.466	1.457	2.262	1.236
ļ	2294	0.713	1.800	1.955	0.663	0.466	1.457	2.262	1.236
	2300	0.713	1.337	2.865	1.515	1.617	1.301	2.098	1.733
	2307	1.779	1.483	2.427	1.764		1.231	1.413	1.000
	2309	1.496	1.483	2.252	1.342		1.278	2.179	4.223
	2313	1.452	2.219	4.257	0.744		1.127	1.588	1.634
	2314	2.044	1.337	2.865	1.515		1.301	2.098	1.733
	2316	1.779	1.337	2.169	1.462		1.784	1.937	2.633
	2327	1.778	1.288	2.075	2.527			3.772	3.393
	2348	2.064	1.200	2.013	2.027		<u> </u>		

Table 11

- 1	2224	2 2 4 2	0.001	0.001	2 225	4 000	4 5000	1 1 1	1 200
- 1	2384	2.340	0.001	0.001	2.927	4.830	1.708	1.651	1.586
- 1	4 2JUT)	4.540	0.001	0.001	4.24	7.000	1.,00	1.051	1.500

Table 12

SEQ ID	1				7			
NO	P268	P278	P295	P339	P341	P356	P360	P392
18	1.000	2.819	1.000		1.238	1.784	0.748	2.486
22	1.194	1.000	1.000		3.006	2.766	1.622	10.061
127	2.953	2.030	8.118	<del></del>	2.854	1.000	1000.000	0.001
139	1000.000	1.332	1.000		1.537	1.000	0.001	0.464
148	2.953	2.030	8.118		2.854	1.000	1000.000	0.001
155	1.187	1.447	1.000		3.621	3.844	1.995	1.313
246	1.422	2.018	2.385	1.218	2.039	3.486	1.636	1.623
272	1.268	1.563	1.870 1.000	2.056	6.240	6.491	2.230	1.427
273	1.000	1000.000			2.209	1000.000	0.001 2.679	
279 298	1.000	1.000	1.000		2.382	3.061	0.001	1.361
329	2.467	2.166	21.707	0.615	1.616	1.000	1.000	1.000
353	1.187	1.447	1.000		3.621	3.844	1.995	1.313
354	2.359	1.552	2.918		4.706	3.623	1.979	1.677
381	2.953	2.030	8.118		2.854	1.000	1000.000	0.001
389	2.467	2.166	21.707	0.615	1.616	1.000	1.000	1.000
405	1.221	1.796	1.995	1.780	1.726	2.970	1.792	1.581
406	1.221	1.796	1.995		1.726	2.970	1.792	1.581
412	2.677	2.809	2.969		2.087	3.804	1.612	1.163
421	2.468	5.262	4.008		4.366	2.078	1.781	1.332
465	1000.000	1.332	1.000		1.537	1.000	0.001	0.464
534	1.221	1.796	1.995	1.780	1.726	2.970	1.792	1.581
538	2.565	1.856	1.000	1.000	2.449	1.000	2.097	2.647
565	1.187	1.447	1.000	1.484	3.621	3.844	1.995	1.313
657	1.369	1.000	1.000	1.679	3.084	2.855	2.104	0.927
670	1.677	2.420	2.263	1.314	1.473	2.523	1.776	2.244
739	1.412	1.431	3.103	1.000	2.847	2.621	1.000	1.117
741	2.240	2.040	1.000		2.450	3.440	2.045	1.998
744	1.837	2.201	2.518		2.248	2.989	1.570	1.409
755	1.000	1.320	0.556		1.321	1.000	1.000	6.185
757	0.713	1000.000	0.632	2.389	0.202	1.000	1.000	0.356
810	1.000	1.320	0.556		1.321	1.000	1.000	6.185
811	1.000	1.320	0.556		1.321	1.000	1.000	6.185 0.734
845 861	2.151	2.384	2.417 9.879		2.327	2.652 0.001	1.000	0.734
915	1.000	2.819	1.000		1.238	1.784	0.748	2.486
954	1.000	1.509		1000.000	2.327	0.001	1.236	0.870
955	1.657	1.732	3.510		4.946	4.071	2.194	1.932
991	1.657	1.732	3.510		4.946	4.071	2.194	1.932
1035	1.677	2.420	2.263	<del></del>	1.473	2.523	1.776	2.244
1049	1.412	1.431	3.103		2.847	2.621	1.000	1.117
1050	1.657	1.732	3.510		4.946	4.071	2.194	1.932
1235	1.187	1.447	1.000		3.621	3.844	1.995	1.313
1292	0.718	1.000	1.000		2.301	1.361	2.161	1.825
1313	1.677	2.420	2.263		1.473	2.523	1.776	2.244

Table 12

1331	0.789	1.609	1.000	0.797	1.000	2.075	2.491	2.505
1334	2.359	1.552	2.918	1.647	4.706	3.623	1.979	1.677
1418	1.000	2.819	1.000	1.589	1.238	1.784	0.748	2.486
1419	1.187	1.447	1.000	1.484	3.621	3.844	1.995	1.313
1420	1.864	1.428	2.631	1.854	3.430	3.182	1.892	1.581
1477	1.864	1.428	2.631	1.854	3.430	3.182	1.892	1.581
1554	2.495	2.090	3.320	1.000	3.907	2.976	1.875	1.000
1579	1.422	2.018	2.385	1.218	2.039	3.486	1.636	1.623
1600	1.187	1.447	1.000	1.484	3.621	3.844	1.995	1.313
1639	1.268	1.563	1.870	2.056	6.240	6.491	2.230	1.427
1657	2.183	2.285	3.554	1.247	2.093	1.840	1.855	1.504
1679	2.495	2.090	3.320	1.000	3.907	2.976	1.875	1.000
1744	2.006	1.696	2.261	1.611	2.154	3.791	1.816	1.356
1847	1.535	2.851	4.154	2.055	6.047	4.103	3.367	2.029
1877	1.187	1.447	1.000	1.484	3.621	3.844	1.995	1.313
1880	2.274	1.266	4.526	2.591	5.409	3.138	2.675	1.391
1889	1.187	1.447	1.000	1.484	3.621	3.844	1.995	1.313
2009	1.000	2.819	1.000	1.589	1.238	1.784	0.748	2.486
2023	1.971	1.699	2.355	1.453	3.122	2.528	1.949	1.326
2029	1.422	2.018	2.385	1.218	2.039	3.486	1.636	1.623
2071	1.677	2.420	2.263	1.314	1.473	2.523	1.776	2.244
2077	1.677	2.420	2.263	1.314	1.473	2.523	1.776	2.244
2103	2.516	0.852	1.775	0.818	4.294	2.281	1.119	0.890
2109	1.677	2.420	2.263	1.314	1.473	2.523	1.776	2.244
2138	1.794	1.486	5.006	0.398	4.768	0.001	2.344	2.434
2143	2.079	1.664	1.000	1.871	2.812	2.693	5.094	1.947
2183	2.325	2.043	2.530	2.411	5.749	5.509	3.490	2.008
2185	1.000	1.320	0.556	1.385	1.321	1.000	1.000	6.185
2190	1.677	2.420	2.263	1.314	1.473	2.523	1.776 1.995	2.244
2200	1.187	1.447	1.000	1.484	3.621 2.538	3.844 1.867	2.312	1.313
2220	2.942	0.729	1.772	0.861	15.794	2.349	1.363	0.808
2224	1.457	1.690	2.551	1.860	4.114	3.548	3.125	0.808
2249	1.187	1.447	1.000	1.484	3.621	3.844	1.995	1.313
2255	1.586	1.943	1.000	0.699	1.593	2.039	1.798	0.774
2265	2.157	1.922	3.895	4.143	2.655	1.914	2.159	3.312
2267	3.442	3.933	5.994	1.448	8.695	7.488	2.687	2.449
2281	2.467	1.000	7.584	1.417	3.693	1.947	1.539	4.429
2294	1.000	2.819	1.000	1.589	1.238	1.784	0.748	2.486
2300	1.000	2.819	1.000	1.589	1.238	1.784	0.748	2.486
2307	1.422	2.018	2.385	1.218	2.039	3.486	1.636	1.623
2309	2.485	2.369	1.000	1.820	3.354	5.046	1.820	0.703
2313	3.203	1.593	4.012	1.593	6.374	6.940	3.158	0.947
2314	1.268	1.563	1.870	2.056	6.240	6.491	2.230	1.427
2316	1.422	2.018	2.385	1.218	2.039	3.486	1.636	1.623
2327	2.439	1.482	2.156	1.390	3.500	3.654	1.655	0.771
2348	2.448	2.617	4.003	1.289	2.940	3.894	2.277	1.202

Table 12

2204	2 220	1 359	0.252	0.202	1 925	0.001	1 1 000	0.714
2384	2.328	1.339	9.253	0.383	1.835	0.001	1.000	I U./141

Table 13

SEQ ID	1			1				
NO	P393	P413	P505	P517	P534	P546	P577	P695
18	1.058	2.471	1.583	1.726	0.506	1.431	2.632	5.930
22	14.260	2.516	1.498	3.747	1.300	5.779	11.202	0.001
127	1.000	0.001	1.000	1.000	0.001	1.000	3.303	1.000
139	1.000	1.000	0.458	1.249	0.001	1000.000	0.702	1.000
148	1.000	0.001	1.000	1.000	0.001	1.000	3.303	1.000
155	1.137	2.268	2.414	1.382	2.107	2.210	2.384	5.256
246	0.741	2.181	2.494	1.504	1.511	1.831	2.064	4.421
272	1.348	2.222	2.506	1.355	1.670	2.535	1.556	8.411
273	1.000	1.000	1000.000	1.477	1.645	1.000	1.389	1.000
279	0.914	1.603	1.936	1.485	2.430	1.999	1.647	4.375
298	1.000	1.000	1000.000	1.477	1.645	1.000	1.389	1.000
329	1.000	1.000	1.436	0.517	1.000	1.469	1.000	1.000
353	1.137	2.268	2.414	1.382	2.107	2.210	2.384	5.256
354	1.224	3.432	2.806	1.328	2.470	2.592	1.929	6.973
381	1.000	0.001	1.000	1.000	0.001	1.000	3.303	1.000
389	1.000	1.000	1.436	0.517	1.000	1.469	1.000	1.000
405	1.241	1.841	1.470	1.000	1.672	2.218	1.649	7.555
406	1.241	1.841	1.470	1.000	1.672	2.218	1.649	7.555
412	1.258	2.153	1.849	1.445	1.000	1.531	1.637	3.302
421	1.000	1.327	2.871	1.116	1.903	2.200	2.644	0.001
465	1.000	1.000	0.458	1.249	0.001	1000.000	0.702	1.000
534	1.241	1.841	1.470	1.000	1.672	2.218	1.649	7.555
538	1.560	1.982 2.268	2.159 2.414	1.278	1.425 2.107	1.204 2.210	3.046 2.384	2.068 5.256
565 657	0.763	1.602	2.797	1.265	2.765	2.236	2.548	5.071
670	1.710	2.337	1.898	0.892	1.347	1.908	1.136	3.404
739	2.102	1.689	4.429	0.830	1.000	1.000	2.108	2.208
741	1.935	1.911	2.812	1.000	1.854	1.793	2.441	0.001
744	1.320	1.404	1.553	1.000	1.957	1.816	2.156	3.745
755	1.219	2.547	1.288	2.539	3.936	3.625	2.363	1.955
757	0.851	0.750	0.815	0.258	0.712	1.229	0.190	1.000
810	1.219	2.547	1.288	2.539	3.936	3.625	2.363	1.955
811	1.219	2.547	1.288	2.539	3.936	3.625	2.363	1.955
845	2.765	1.000	2.202	0.472	0.490	1.417	0.725	0.001
861	1.000	1.000	1.000	1.000	1.000	1.530	0.769	1.000
915	1.058	2.471	1.583	1.726	0.506	1.431	2.632	5.930
954	1.000	1.000	1.000	1.000	1.000	1.530	0.769	1.000
955	1.322	2.608	1.910	1.199	1.635	1.893	1.473	5.842
991	1.322	2.608	1.910	1.199	1.635	1.893	1.473	5.842
1035	1.710	2.337	1.898	0.892	1.347	1.908	1.136	3.404
1049	2.102	1.689	4.429	0.830	1.000	1.000	2.108	2.208
1050	1.322	2.608	1.910	1.199	1.635	1.893	1.473	5.842
1235	1.137	2.268	2.414	1.382	2.107	2.210	2.384	5.256
1292	1.000	1.518	1.980	1.518	2.526	1.588	1.865	2.251
1313	1.710	2.337	1.898	0.892	1.347	1.908	1.136	3.404

Table 13

1331	0.743	2.126	1.613	1.177	2.128	1.000	1.951	6.931
1334	1.224	3.432	2.806	1.328	2.470	2.592	1.929	6.973
1418	1.058	2.471	1.583	1.726	0.506		2.632	5.930
1419	1.137	2.268	2.414	1.382	2.107	2.210	2.384	5.256
1420	1.205	3.301	2.749	1.256	2.474		1.826	8.108
1477	1.205	3.301	2.749	1.256	2.474		1.826	8.108
1554	1.000	1.793	2.719	1.679	1.000		2.076	0.001
1579	0.741	2.181	2.494	1.504	1.511	1.831	2.064	4.421
1600	1.137	2.268	2.414	1.382	2.107		2.384	5.256
1639	1.348	2.222	2.506	1.355	1.670		1.556	8.411
1657	2.809	1.534	1.366		2.545	1.964	1.506	0.001
1679	1.000	1.793	2.719	1.679	1.000	1.549	2.076	0.001
1744	1.249	2.009	1.832	1.488	1.379	1.975	2.128	13.930
1847	1.781	2.929	2.183	2.759	3.853	3.092	2.051	7.549
1877	1.137	2.268	2.414	1.382	2.107	2.210	2.384	5.256
1880	1.000	3.187	2.564	0.756	1.226		3.201	16.724
1889	1.137	2.268	2.414	1.382	2.107	2.210	2.384	5.256
2009	1.058	2.471	1.583	1.726	0.506		2.632	5.930
2023	1.952	1.472	1.917	1.516	2.305	2.677	2.620	2.660
2029	0.741	2.181	2,494		1.511	1.831	2.064	4.421
2071	1.710	2.337	1.898	0.892	1.347	1.908	1.136	3.404
2077	1.710	2.337	1.898	0.892	1.347	1.908	1.136 1.239	3.404 0.001
2103	0.537	1.790	0.727	0.750 0.892	0.329 1.347	1.100 1.908	1.136	3.404
2109	1.710 0.852	2.337 1.789	1.898 3.765	0.892			1.852	0.001
2143	2.044	17.760	4.034		0.026		2.394	42.662
2183	1.088	5.833	3.519	1.572	2.641	4.011	1.695	7.783
2185	1.219	2.547	1.288	2.539			2.363	1.955
2190	1.710	2.337	1.898	0.892	1.347	1.908	1.136	
2200	1.137	2.268	2.414	1.382	2.107	2.210		5.256
2206	1.000	3.033	1.912	1.699		2.780	2.155	2.518
2220	0.337	2.339	0.768	0.563	0.359	1.242	1.492	1.000
2224	1.000	2.266		1.000		2.620	1.718	14.145
2249		2.268			1			
2255	1.243	1.766						
2265	5.268	1.518	2.253	3.678	0.766	1.565	1.000	1.853
2267	0.815	2.497	3.234	2.275	2.344	3.596	5.023	12.124
2281	1.128	0.885	1.237	1.434	3.327	3.206	1.355	0.001
2294	1.058	2.471	1.583	1.726	0.506	1.431	2.632	5.930
2300	1.058	2.471	1.583	1.726	0.506	1.431	2.632	5.930
2307	0.741	2.181	2.494	1.504	1.511	1.831	2.064	4.421
2309	1.240	2.239	2.841	1.000	2.270	2.614	0.583	5.244
2313	0.633	2.821	2.976	1.253	1.675	3.657	2.284	8.587
2314	1.348	2.222	2.506	1.355	1.670	2.535	1.556	
2316	0.741	2.181	2.494		<del></del>		2.064	
2327	1.000	1.801	1.978				2.276	
2348	0.790	3.524	3.377	2.062	2.123	1.959	1.626	1.000

Table 13

į	2204	0.001	1 246	1.021	1 000	1 (46	1.944	1.540	1.000
	2384	0.001	1.346	1.831	1.000	1.646	1.944	1.549	1.000

Table 14

SEQ ID			F		
NO	P784	P786	P791	P888	P889
18	1.000	1.000	4.202	1.464	2.147
22	1.000	1.276	14.034	4.139	3.640
127	1.708	2.247	1.000	0.441	0.001
139	1.391	1.857	1.000	0.402	1.000
148	1.708	2.247	1.000	0.441	0.001
155	1.328	1.421	2.456	1.910	2.069
246	1.243	1.679	2.228	2.333	1.774
272	0.819	1.632	2.808	5.465	2.307
273	1000.000	0.758	1.000	1.000	1.000
279	1.000	1.000	1.834	2.776	1.636
298	1000.000	0.758	1.000	1.000	1.000
329	1.000	1.000	1.000	0.642	1.000
353	1.328	1.421	2.456	1.910	2.069
354	1.000	1.416	2.862	2.690	1.645
381	1.708	2.247	1.000	0.441	0.001
389	1.000	1.000	1.000	0.642	1.000
405	1.000	1.821	1.628	2.276	2.501
406	1.000	1.821	1.628	2.276	2.501
412	1.000	1.888	1.915	2.276	1.481
421	3.336	1.677	2.208	1.000	1.976
465	1.391	1.857	1.000	0.402	1.000
534	1.000	1.821	1.628	2.276	2.501
538	1.000	1.629	2.152	1.000	1.792
565	1.328	1.421	2.456	1.910	2.069
657	1.000	1.997	2.083	3.178	3.444
670	1.000	1.780	1.000	2.177	2.258
739	1.356	0.696	1.000	1.000	1.463
741	2.324	1.000	2.379	1.407	2.833
744 755	2.137	1.934	2.482	2.035	3.980
757	0.796	1.000	1.737	1.000	2.218
	2.531	3.138	0.395	1.000	1.000
810 811	0.796	1.000	1.737	1.000	2.218
845	0.796 1000.000	1.000	1.737	1.000	2.218
861	3.031	1.984	1000.000	1.374	1.000
915	1.000	1.000	1.000	1.000	1.000
954	3.031		4.202	1.464	2.147
955	0.876	1.000	1.000	1.000	1.000
991	0.876	1.781 1.781	2.424	4.143	1.977
1035	1.000		2.424	4.143	1.977
1049	1.356	1.780 0.696	1.000	2.177	2.258
1050	0.876	1.781	1.000	1.000	1.463
1235	1.328	1.421	2.424	4.143	1.977
1292	1.000	1.000	2.456	1.910	2.069
1313	1.000	1.780	1.992	2.144	1.615
	1.000	1.700	1.000	2.177	2.258

Table 14

1331	1.290	1.000	1.000	1.995	2.203
1334	1.000	1.416	2.862	2.690	1.645
1418	1.000	1.000	4.202	1.464	2.147
1419	1.328	1.421	2.456	1.910	2.069
1420	0.816	1.000	2.196	2.446	1.518
1477	0.816	1.000	2.196	2.446	1.518
1554	1.585	1.889	2.178	1.806	1.867
1579	1.243	1.679	2.228	2.333	1.774
1600	1.328	1.421	2.456	1.910	2.069
1639	0.819	1.632	2.808	5.465	2.307
1657	2.810	2.638	1.976	1.491	2.955
1679	1.585	1.889	2.178	1.806	1.867
1744	1.253	1.994	1.874	3.193	2.663
1847	1.559	2.762	5.043	4.135	3.753
1877	1.328	1.421	2.456	1.910	2.069
1880	1.306	1.940	2.293	3.897	1.624
1889	1.328	1.421	2.456	1.910	2.069
2009	1.000	1.000	4.202	1.464	2.147
2023	1.511	1.357	1.632	1.891	1.895
2029	1.243	1.679	2.228	2.333-	1.774
2071	1.000	1.780	1.000	2.177	2.258
2077	1.000	1.780	1.000	2.177	2.258
2103	0.573	2.678	1.000	2.507	3.278
2109	1.000	1.780	1.000	2.177	2.258
2138	7.866	1.000	1000.000	1.719	1.000
2143	2.625	2.744	4.155	2.105	4.438
2183	1.000	2.139	3.014	3.159	3.381
2185	0.796	1.000	1.737	1.000	2.218
2190	1.000	1.780	1.000	2.177	2.258
2200	1.328	1.421	2.456	1.910	2.069
2206	1.489	2.750	2.910	5.049	4.006
2220	0.419	3.014	0.575	2.397	3.558
2224	1.000	1.815	2.513	3.487	2.180
2249	1.328	1.421	2.456	1.910	2.069
2255	1.000	1.493	2.186	1.000	2.222
2265	1.267	3.638	1.623	5.889	3.339
2267	1.746	2.363	5.515	2.674	3.637
2281	2.399	3.587	3.625	2.567	2.417
2294	1.000	1.000	4.202	1.464	2.147
2300	1.000	1.000	4.202	1.464	2.147
2307	1.243	1.679	2.228	2.333	1.774
2309	0.397	1.000	1.472	5.315	2.250
2313	1.000	1.939	2.505	4.525	2.674
2314	0.819	1.632	2.808	5.465	2.307
2316	1.243	1.679	2.228	2.333	1.774
2327	1.295	1.658	2.836	2.766	2.873
2348	2.167	2.157	3.410	2.828	3.794

Table 14

2384	1.352	1 0001	0.707	0.70-	
1 43041	1.3321	1.000	9.797	0.522	1 1 1 1 1 1 1 1
	~	1.000	2.121	0.583	1.000

Table 1

	able 1				
SEQ ID			ORIENTATION	CLONE ID	LIBRARY
1	734646	RTA22200010F.k.10.1.P.Seq	F	M00056481:62	
2	400221	RTA22200001F.a.17.1.P.Seq	F	M00042528:611	
3	205329	RTA22200006F.d.09.2.P.Seq	F F	M00056020:410	
4	446680	RTA22200001F.f.07.1.P.Seq	F.	M00042693:54	
5	1261	RTA22200021F.j.18.3.P.Seq	F	M00054812:15	
6	400258	RTA22200011F.k.23.1.P.Seq	F	M00054612:15	; CH16COP
7	450559	RTA22200005F.e.21.1.P.Seq	F	M00055882:16	
8	450959	RTA22200012F.e.11.1.P.Seq	~~~~ <b>~</b>	WASTER THE HOME MANAGEMENT AND ASSESSMENT OF THE PARTY OF	CH15CON
9	451794	RTA2220007F.I.16.1.P.Seq	F	M00056703:46	CH16COP
10	415058	RTA222000071.1.10.1.F.Seq	F	M00056247:76	CH15CON
11	31506	RTA22200020F.d.11.1.P.Seq		M00054591:87	CH17COHLV
12	417155	DTA22200012F.b.06.1.P.Seq	<u> </u>	M00056670:111	restaurant actus autorium autorium a const at man an an
13		RTA22200002F.f.10.1.P.Seq	F	M00055466:28	was about the contract of the
	448925	RTA22200019F.e.21.1.P.Seq	F	M00043507:45	CH17COHLV
14	11329	RTA22200006F.d.10.2.P.Seq	F	M00056020:47	CH15CON
15	650422	RTA22200001F.n.14.1.P.Seq	F	M00042911:83	CH15CON
16	6863	RTA22200229F.f.13.1.P.Seq	F	M00006967:25	CH02COH
17	449690	RTA22200002F.g.18.1.P.Seq	F	M00055495:45	CH15CON
18	724616	RTA22200016F.j.23.1.P.Seq	F	M00057236:86	CH16COP
19	549722	RTA22200025F.m.01.2.P.Seq	F	M00055383:24	CH17COHLV
20	549722	RTA22200025F.I.24.1.P.Seq	F	M00055383:24	CH17COHLV
21	448110	RTA22200018F.m.04.1.P.Seq	F	M00043354:31	
22	515631	RTA22200010F.j.14.1.P.Seg	F	M00056434:38	CH16COP
23	11881	RTA22200233F.k.04.1.P.Seq	F	M00008099:78	CH03MAH
24	650856	RTA22200012F.n.24.1.P.Seq	F	M00056772:14	CH16COP
25	449701	RTA22200012F.f.21.1.P.Seq		M00056710:89	CH16COP
26	651073	RTA22200007F.I.06.1.P.Seg	~	M00056243:710	
27	10340	RTA22200234F.b.07.1.P.Seq	F	M00030243.710 M00022189:23	
28	648310	RTA22200007F.m.04.1.P.Seq	F	NONETHINGS OF THE PROPERTY OF	CH03MAH
29	730336	RTA22200013F.I.02.1.P.Seq	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M00056252:88	CH15CON
30	3060	RTA22200018F.b.10.1.P.Seq	F	M00056879:811	CH16COP
31	453016	RTA22200010F.I.06.1.P.Seq	······	M00042444:88	CH17COHLV
32	508931	RTA222000101.i.00.1.F.Seq	men entre eller monomonere el energialistica de la	M00056485:212	CH16COP
33	185461	RTA22200024F.i. 13.1.P.Seq	F	M00055209:410	personal and an accompany of the second and the second accompany of the second
34	452530		<u> </u>	M00026975:23	CH04MAL
35	448925	RTA22200015F.n.11.1.P.Seq		M00057131:21	CH16COP
36	CONTRACTOR A MONTH TO THE MANAGEMENT	RTA22200026F.d.02.1.P.Seq	entering and recognition control and anterior and include control trighter	M00055419:71	CH17COHLV
37	·····	RTA22200005F.m.06.1.P.Seq	***************************************	M00055945:811	CH15CON
·····	6545	RTA22200241F.d.23.1.P.Seq	arania and a sanda and a s	M00026879:410	CH04MAL
38	449891	RTA22200001F.b.23.1.P.Seq	***************************************	M00042540:85	CH15CON
39	4045	RTA22200227F.n.06.1.P.Seq		M00006740:71	CH02COH
40		RTA22200002F.b.23.1.P.Seq		M00055438:810	CH15CON
41		RTA22200001F.n.10.1.P.Seq ,		M00042909:74	CH15CON
42	650493	RTA22200005F.n.03.1.P.Seq	F	M00055959:112	CH15CON
43		RTA22200007F.k.04.1.P.Seq		M00056232:712	CH15CON
44	452212	RTA22200021F.k.21.3.P.Seq	a conserve reconserverse reconservers as according ,	ACCOMPANIES DECIMAL NEW TOTAL CONTROL OF THE STREET SAN	CH17COHLV
45		RTA22200010F.n.09.1.P.Seq	***************************************	M00056505:82	CH16COP
46		RTA22200003F.m.24.1.P.Seq		M00055709:79	CH15CON
47		RTA22200013F.f.14.1.P.Seq	remenenament in interest in the contract of th	M00056839:72	CH16COP
48		RTA22200003F.o.06.1.P.Seq	MARKET AND	M00055723:28	CH15CON
49		RTA22200001F.e.15.1.P.Seq	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M00033723.28	CH15CON
50	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	RTA22200012F.o.07.1.P.Seq		M00042370.82   M00056774:12	······································
51	······································	RTA22200010F.i.17.1.P.Seq	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	······································	CH16COP
52	accompanies and a second	RTA22200018F.k.24.1.P.Seq		M00056424:110	CH16COP
53		RTA22200010F.R.24.1.P.Seq	and the second s		CH17COHLV
54		RTA222000011.111.13.1.P.Seq		M00042902:34	CH15CON
	11020	1.1742200230F.y.12.1.P.Seq	F N	100007151:211	CH02COH

Table 1

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57	446789	RTA22200001F.c.12.1.P.Seq	F	M00042547:62	CH15CON
58	644884	RTA22200007F.k.04.2.P.Seq	F	M00056232:712	
59	9029	RTA22200222F.g.08.1.P.Seq	F	M00003792:111	·2·····
60	419255	RTA22200007F.I.14.2.P.Seq	F	M00056246:77	CH15CON
61	4309	RTA22200225F.j.15.1.P.Seq	F	M00005491:23	CH02COH
62	554069	RTA22200022F.p.16.1.P.Seq	F	M00055008:28	CH17COHLV
63	4330	RTA22200227F.i.16.1.P.Seq	F	M00006686:27	CH02COH
64	644903	RTA22200015F.i.22.1.P.Seq	F	M00057103:511	CH16COP
65	549395	RTA22200024F.d.04.1.P.Seq	F	M00055163:32	CH17COHLV
66	4974	RTA22200225F.k.17.1.P.Seq	F	M00005500:53	CH02COH
67	447466	RTA22200006F.h.03.2.P.Seq	F	M00005500:55 M00056053:19	CH15CON
68	645073	RTA22200004F.o.13.1.P.Seq	F	M00055829:79	CH15CON
69	447978	RTA22200025F.i.14.1.P.Seq	F	M00055829.79	CH17COHLV
70	607430	RTA222000237 .1.14.1.F.3eq	F	M00055841:45	CH17CORLV
71	556198	RTA22200004F.h.02.2.P.Seq	F		denne de la companya
72	450323	RTA22200014F.II.02.2.F.Seq	F	M00056972:65	CH16COP
73	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~			M00043352:59	CH17COHLV
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75 76	446673	RTA22200001F.a.14.1.P.Seq	<u> </u>	M00042525:51	CH15CON
76	456026	RTA22200004F.m.04.1.P.Seq	<u> </u>	M00055817:38	CH15CON
77	449142	RTA22200009F.c.13.2.P.Seq	<u> </u>	M00042513:112	CH16COP
78	5830	RTA22200010F.c.23.1.P.Seq	F	M00056370:72	CH16COP
79	554109	RTA22200026F.g.07.1.P.Seq	F	M00055484:77	CH17COHLV
80	595506	RTA22200010F.I.16.1.P.Seq	F	M00056491:78	CH16COP
81	453981	RTA22200010F.p.11.1.P.Seq	F	M00056519:81	CH16COP
82	642461	RTA22200016F.j.11.1.P.Seq	F	M00057233:54	CH16COP
83	556198	RTA22200014F.h.02.1.P.Seq	F	M00056972:65	CH16COP
84	2082	RTA22200009F.g.21.1.P.Seq	F	M00042801:26	CH16COP
85	549435	RTA22200007F.j.06.1.P.Seq	F	M00056223:73	CH15CON
86	2286	RTA22200230F.j.03.1.P.Seq	F	M00007177:511	CH02COH
87	2737	RTA22200023F.d.17.1.P.Seq	F	M00055039:52	CH17COHLV
88	728115	RTA22200013F.f.13.1.P.Seq	F	M00056839:71	CH16COP
89	650856	RTA22200012F.o.01.1.P.Seq	F	M00056772:14	CH16COP
90	650476	RTA22200005F.g.21.1.P.Seq	F	M00055891:14	CH15CON
91	535208	RTA22200005F.b.02.1.P.Seq	F	M00055852:17	CH15CON
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93	447978	RTA22200009F.g.19.1.P.Seq	F :	M00042800:13	CH16COP
94	729483	RTA22200012F.i.12.1.P.Seq	F	M00056728:45	CH16COP
95	12018	RTA22200249F.e.19.1.P.Seq	F	M00027641:11	CH04MAL
96	4747	RTA22200227F.d.18.1.P.Seq	xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx	M00006630:311	CH02COH
97	4747	RTA22200225F.a.23.1.P.Seq	·····	M00005415:612	CH02COH
98	185577	RTA22200240F.i.02.1.P.Seq	F	M00023409:78	CH04MAL
99	4126	RTA22200231F.m.17.1.P.Seq	F	M00007990:43	CH03MAH
100	11456	RTA22200226F.I.08.1.P.Seq	F	M00005765:67	CH02COH
101	729851	RTA22200010F.o.03.1.P.Seq	**************************************	M00056508:210	CH16COP
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103	2490	RTA22200237F.e.17.1.P.Seq	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M00022720:111	CH03MAH
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105	11881	RTA22200237F.i.01.1.P.Seq	F	M00034779:07	CH03MAH
106	724296	RTA22200014F.o.10.1.P.Seq	F	M00057025:18	CH16COP
107	726173	RTA22200014F.o.01.1.P.Seq	F	M00057023:18	CH16COP
108	2423	RTA22200236F.h.20.1.P.Seq	F	M00037023.69 M00022641:62	CH03MAH
		ELLOUZOUI .II.ZU. I.IOEY		1V100022041.02	OLIOSIVIAL

Table 1 **CLONE ID** LIBRARY **ORIENTATION** SEQ NAME SEQ ID CLUSTER CH16COP M00057005:31 556250 RTA22200014F.l.09.1.P.Seq 109 F M00055851:612 CH15CON RTA22200005F.b.01.1.P.Seq 110 643594 CH03MAH F 111 11881 RTA22200237F.h.24.1.P.Seq M00022750:17 RTA22200237F.j.11.1.P.Seq F M00022791:611 CH03MAH 112 7436 F M00022415:26 CH03MAH RTA22200235F.a.19.1.P.Seq 113 2110 F M00022457:75 CH03MAH RTA22200235F.e.13.1.P.Seq 114 10340 F M00055851:612 CH15CON 643594 RTA22200005F.a.24.1.P.Seq 115 CH15CON F M00055872:412 RTA22200005F.d.11.1.P.Seq 116 447035 CH15CON RTA22200007F.c.21.1.P.Seq F M00056160:18 117 402707 CH16COP M00056952:84 F 118 645799 RTA22200014F.d.22.1.P.Seq CH16COP RTA22200014F.c.05.1.P.Seq F M00056939:22 119 171511 CH16COP RTA22200014F.h.15.2.P.Seq F M00056976:610 120 451607 RTA22200242F.k.08.1.P.Seq F M00027039:59 CH04MAL 3138 121 F M00006987:711 CH02COH RTA22200229F.h.11.1.P.Seq 122 2988 F CH16COP M00056969:21 RTA22200014F.g.22.1.P.Seq 123 447326 CH16COP F M00056961:712 124 561734 RTA22200014F.f.10.1.P.Seq CH16COP M00042432:810 F 125 454999 RTA22200009F.a.13.2.P.Seq CH04MAL F M00027028:37 185652 RTA22200242F.i.12.1.P.Seq 126 CH03MAH M00021947:36 RTA22200232F.d.17.1.P.Seq F 127 6725 F M00056864:89 CH16COP 726644 RTA22200013F.i.19.1.P.Seq 128 F M00006745:12 CH02COH RTA22200227F.n.21.1.P.Seq 129 11012 F M00056860:612 CH16COP RTA22200013F.i.03.1.P.Seq 130 726377 F M00056886:311 CH16COP RTA22200013F.I.19.1.P.Seq 131 735326 F CH16COP M00056884:36 RTA22200013F.I.13.1.P.Seq 132 650845 CH01COH F M00004101:81 133 9048 RTA22200222F.I.18.1.P.Seq CH16COP F M00056674:84 134 732254 RTA22200012F.b.15.1.P.Seq CH16COP RTA22200013F.i.16.1.P.Seq F M00056863:53 452052 135 CH16COP F M00056735:28 554079 RTA22200012F.j.17.1.P.Seq 136 F CH01COH RTA22200222F.i.05.1.P.Seq M00003948:212 9049 137 CH04MAL F M00027222:39 RTA22200244F.n.16.1.P.Seq 1307 138 CH04MAL F M00027016:76 RTA22200242F.g.11.1.P.Seq 139730 139 F M00026899:711 CH04MAL RTA22200241F.f.24.1.P.Seq 7750 140 F M00006761:49 CH02COH RTA22200227F.p.20.1.P.Seq 141 8050 CH16COP F M00056879:55 RTA22200013F.k.24.1.P.Seq 142 725222 CH03MAH F M00022516:59 143 3275 RTA22200235F.j.22.2.P.Seq CH03MAH F M00022430:44 144 7424 RTA22200235F.c.12.1.P.Seq F M00026866:88 CH04MAL RTA22200241F.c.05.1.P.Seq 145 8953 F M00027088:86 CH04MAL RTA22200243F.c.04.1.P.Seq 8966 146 CH16COP F M00056866:55 RTA22200013F.i.22.1.P.Seq 530883 147 CH03MAH F M00022973:78 6725 RTA22200238F.I.09.1.P.Seq 148 M00004167:411 CH01COH F 4439 RTA22200222F.m.24.1.P.Seq 149 M00056712:17 CH16COP RTA22200012F.g.07.1.P.Seq F 150 648472 CH16COP M00056618:22 F 735346 RTA22200011F.I.03.1.P.Seq 151 F M00056600:87 CH16COP RTA22200011F.j.05.1.P.Seq 152 732121 F CH15CON M00055900:25 153 650337 RTA22200005F.h.15.1.P.Seq CH15CON F M00055981:17 RTA22200005F.p.05.1.P.Seq 154 533588 CH15CON F M00056290:82 RTA22200007F.p.20.1.P.Seq 155 649667 CH16COP F M00057145:45 RTA22200015F.p.07.1.P.Seq 394436 156 F M00056210:53 CH15CON RTA22200007F.h.13.1.P.Seq 649354 157 M00023347:312 CH04MAL F RTA22200240F.e.10.1.P.Seq 158 2022 CH15CON F M00055703:26 RTA22200003F.m.08.1.P.Seq 561359 159 F M00005520:512 CH02COH RTA22200225F.m.20.1.P.Seq 160 7607 F M00027006:81 CH04MAL RTA22200242F.f.06.1.P.Seq 7750 161 F M00056729:812 CH16COP 410554 RTA22200012F.i.21.1.P.Seq 162

Table 1 Page 3 of 45

Table 1

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163	2315	RTA22200230F.e.24.1.P.Seq	F	M00007135:211	CH02COH
164	561734	RTA22200014F.f.10.2.P.Seq	F	M00056961:712	CH16COP
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167	7082	RTA22200235F.p.01.2.P.Seq	F	M00022565:15	CH03MAH
168	2315	RTA22200230F.f.01.1.P.Seq	F	M00007135:211	CH02COH
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170	6482	RTA22200230F.a.10.1.P.Seq	F F	M00007096:51	CH02COH
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172	453846	RTA22200012F.b.14.1.P.Seq	F	M00056674:55	CH16COP
173	650820	RTA22200011F.p.05.1.P.Seq	F	M00056656:83	CH16COP
174	642906	RTA22200005F.g.06.1.P.Seq	F	M00055887:36	CH15CON
175	448805	RTA22200005F.i.23.1.P.Seq	F	M00055007:35	CH15CON
176	649667	RTA22200005F.k.18.2.P.Seq	F	M00055912:55	CH15CON
177	735786	RTA22200000F.K.18.2.F.Seq	F	M00056758:35	CH16COP
\$ management and the second		······································	F	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	CH16COP
178	121457	RTA22200012F.p.18.1.P.Seq	3	M00056785:68	}
179	372960	RTA22200012F.m.06.1.P.Seq	5	M00056756:28	CH16COP
180	120049	RTA22200012F.j.10.1.P.Seq	<u> </u>	M00056733:49	CH16COP
181	648996	RTA22200006F.p.20.2.P.Seq	<u>F</u>	M00056136:211	CH15CON
182	3765	RTA22200226F.a.16.1.P.Seq	F	M00005589:67	CH02COH
183	462642	RTA22200008F.e.08.1.P.Seq	F	M00056342:75	CH15CON
184	727181	RTA22200016F.f.18.1.P.Seq	F	M00057208:12	CH16COP
185	649259	RTA22200006F.k.06.2.P.Seq	F	M00056079:412	CH15CON
186	649717	RTA22200007F.e.08.1.P.Seq	F	M00056180:89	CH15CON
187	736860	RTA22200009F.p.24.1.P.Seq	F	M00056351:46	CH16COP
188	729175	RTA22200012F.o.05.1.P.Seq	F	M00056773:811	CH16COP
189	642906	RTA22200005F.f.14.1.P.Seq	F	M00055884:15	CH15CON
190	4420	RTA22200232F.I.13.1.P.Seq	F	M00022123:45	CH03MAH
191	2420	RTA22200226F.c.12.1.P.Seq	F	M00005619:19	CH02COH
192	648109	RTA22200015F.o.03.1.P.Seq	F	M00057135:84	CH16COP
193	2334	RTA22200011F.p.20.1.P.Seq	F	M00056661:19	CH16COP
194	639705	RTA22200003F.o.14.1.P.Seq	F	M00055724:49	CH15CON
195	551907	RTA22200003F.n.12.1.P.Seq	F	M00055717:64	CH15CON
196	561382	RTA22200003F.m.20.1.P.Seq	F	M00055706:71	CH15CON
197	595506	RTA22200022F.a.01.1.P.Seq	F	M00054866:77	CH17COHLV
198	499424	RTA22200013F.f.16.1.P.Seq	F	M00056839:61	CH16COP
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200	734370	RTA22200013F.g.21.1.P.Seq	F	M00056848:37	CH16COP
201	779	RTA22200230F.c.07.1.P.Seq	F	M00007112:112	CH02COH
202	649143	RTA22200007F.o.24.1.P.Seq	F	M00056283:52	CH15CON
203	489	RTA22200012F.m.16.1.P.Seq	F	M00056759:611	CH16COP
204	2994	RTA22200229F.k.24.1.P.Seq	F	M000007028:34	CH02COH
205	2994	RTA22200229F.I.01.1.P.Seq	F	M00007028:34	CH02COH
206	11147	RTA22200223F.i.01.7.F.Seq	F	M00007020:34 M00006664:29	CH02COH
207	549395	RTA222002277 .g. 12.1.7 .Seq	F	M000054745:13	CH17COHLV
208	559806	RTA22200021F.d.19.2.F.Seq	F	M00054743.13	CH17COHLV
208	452238	RTA22200022F.II.11.1.P.Seq	F	M00042838:511	CH1/COHLV
Samuel Marie Commence	<del>ar restaura sa restaura sa</del>		F	***************************************	
210	225914	RTA22200023F.j.18.1.P.Seq		M00055075:85	CH17COHLV
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213	557401	RTA22200023F.g.22.1.P.Seq	<u> </u>	M00055056:26	CH17COHLV
214	455155	RTA22200003F.f.04.1.P.Seq	F	M00055633:711	CH15CON
215	551117	RTA22200023F.p.03.1.P.Seq	F	M00055131:210	(antino comence menor menor con concerno, menor menor
216	7659	RTA22200231F.g.16.1.P.Seq	F	M00007964:71	CH03MAH

Table 1

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217	729295	RTA22200017F.b.14.1.P.Seq	F	M00057306:87	CH16COP
218	450429	RTA22200007F.f.20.1.P.Seq	F	M00056195:38	CH15CON
219	450148	RTA22200022F.f.21.1.P.Seq	F	M00054927:58	CH17COHLV
220	380412	RTA22200006F.c.23.2.P.Seq	F	M00056016:46	CH15CON
221	446614	RTA22200001F.e.01.1.P.Seq	F	M00042563:52	CH15CON
222	555911	RTA22200023F.f.21.1.P.Seq	F	M00055050:74	CH17COHLV
223	450828	RTA22200022F.g.23.1.P.Seq	F	M00054935:74	CH17COHLV
224	28	RTA22200006F.a.17.2.P.Seq	F	M00055999:710	CH15CON
225	446450	RTA22200001F.h.15.1.P.Seg	F	M00042717:44	CH15CON
226	452026	RTA22200010F.g.18.1.P.Seq	F	M00056411:54	CH16COP
227	643594	RTA22200006F.a.18.2.P.Seg	F	M00055999:76	CH15CON
228	1905	RTA22200012F.o.14.1.P.Seq	F	M00056775:38	CH16COP
229	651073	RTA22200007F.I.06.2.P.Seq	F	M00056243:710	CH15CON
230	553705	RTA22200006F.a.23.2.P.Seq	F	M00056001:27	CH15CON
231	521840	RTA22200004F.j.15.1.P.Seq	F	M00055802:84	CH15CON
232	648689	RTA22200006F.o.07.2.P.Seq	F	M00056111:82	CH15CON
233	447858	RTA22200022F.d.10.1.P.Seq	F	M00054895:49	CH17COHLV
234	556198	RTA22200010F.d.10.1.P.Seq	F	M00056374:82	CH16COP
235	394436	RTA22200003F.i.09.1.P.Seq	F	M00055574:02 M00055662:14	CH15CON
236	639651	RTA22200003F.g.12.1.P.Seq	F	M00055647:24	CH15CON
237	499424	RTA222000031.g.12.1.F.Seq	F	M00055047:24 M00056382:82	CH16COP
238	468109	RTA22200010F.p.12.1.P.Seq	F	M00054915:57	CH15CON
239	185701	RTA22200248F.g.23.1.P.Seq	F	M00034913.37	CH04MAL
240	451811	RTA22200006F.h.04.2.P.Seq	F	M00056053:412	CH04MAL CH15CON
241	730670	RTA22200000F.m.14.1.P.Seq	F	M00030033.412	CH16COP
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245	414739	RTA222000011.11.23.11.F.Seq	F	M00054945:77	CH17COHLV
246	641124	RTA22200022F.i.10.11F.Seq	F	M00055805:37	CH15CON
247	555702	RTA222000022F.j.10.1.P.Seq	F	M00053803:57	CH17COHLV
248	549435	RTA22200019F.p.14.1.P.Seq	F	M00054561:56	CH17COHLV
249	643954	RTA222000191.p.14.1.F.Seq	F	M00055451:711	CH17CORLV
250	5984	RTA22200026F.d.05.1.P.Seq	F	M00055420:56	CH17COHLV
251	560526	RTA22200022F.f.10.1.P.Seq	F	M00053420:36	CH17COHLV
252	411113	RTA22200023F.a.16.1.P.Seq	F	M00055017:111	CH17COHLV
253	7607	RTA222000251.a.10.1.F.Seq	F	M00055744:69	CH15CON
254	559409	RTA22200019F.o.07.1.P.Seq		M00054551:73	CH17COHLV
255	650053	RTA222000191.0.07.1.1 .Seq	F	M00055560:62	
256	448511	RTA22200002F.f.03.1.P.Seq	F	M00033300.02 M00042777:46	CH15CON CH16COP
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259	431601	RTA22200006F.b.14.2.P.Seq RTA22200004F.d.07.1.P.Seq	######################################	M00056005:34	CH15CON
260	421431	000000 to 1 con 100000 to 100000 to 100000 con 1000000 to 1000000000000000000000000000	F F	M00055763:56	CH15CON
261	····	RTA22200001F.g.16.1.P.Seq	การ ก็สาราบาน เพราะ (การาบาน เมษายน ค.ศ. ค.ศ. เมษายน ค.ศ. ค.ศ. ค.ศ. ค.ศ. ค.ศ. ค.ศ. ค.ศ. ค.ศ	M00042704:52	CH15CON
262	284586	RTA22200002F.d.17.1.P.Seq	F	M00055451:67	CH17COHIV
francourant for	556198	RTA22200022F.l.06.1.P.Seq	F	M00054963:84	CH17COHLV
263	431601	RTA22200004F.j.04.1.P.Seq	F	M00055800:48	CH15CON
264 265	449891	RTA22200002F.n.21.1.P.Seq	F	M00055548:64	CH15CON
266	556561	RTA22200022F.p.01.1.P.Seq		M00054997:212	т жүнинин күрүүчү жүрүнү жүрүү жүрүү
(marana marana and a	554188	RTA22200022F.o.20.1.P.Seq	<u> </u>	M00054996:39	****
267	3247	RTA22200006F.a.22.2.P.Seq	F	M00056001:26	CH15CON
268 269	546705	RTA22200022F.k.20.1.P.Seq		M00054959:311	·
209	560984	RTA22200022F.p.02.1.P.Seq		M00054997:83	NOT TAKEN OF F. ESPECIAL APPEALANCE AND AND RELIEF
210	455820	RTA22200006F.g.03.2.P.Seq	F	M00056045:81	CH15CON

Table 1

	ble 1				
SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	ALLEGAN PARTICION CONTRACTOR MODELLOS ANTICOS EL CARROL MODELLOS ESPACIAS.	LIBRARY
271	643129	RTA22200006F.n.12.2.P.Seq	F	M00056103:412	CH15CON
272	454653	RTA22200023F.o.20.1.P.Seq	F	M00055128:210	CH17COHLV
273	456549	RTA22200002F.k.01.1.P.Seq	F	M00055522:32	CH15CON
274	454806	RTA22200004F.n.04.1.P.Seq	F	M00055822:84	CH15CON
275	724296	RTA22200014F.o.10.2.P.Seq	F	M00057025:18	CH16COP
276	559280	RTA22200015F.j.19.1.P.Seq	F	M00057106:26	CH16COP
277	171511	RTA22200014F.c.05.2.P.Seq	F	M00056939:22	CH16COP
278	644242	RTA22200002F.o.14.1.P.Seq	F	M00055553:84	CH15CON
279	734370	RTA22200016F.j.07.1.P.Seq	F	M00057232:46	CH16COP
280	639459	RTA22200002F.i.18.1.P.Seq	F	M00055512:76	CH15CON
281	641679	RTA22200003F.d.20.1.P.Seq	F	M00055613:52	CH15CON
282	644611	RTA22200002F.i.10.1.P.Seq	F	M00055509:89	CH15CON
283	550038	RTA22200021F.i.16.3.P.Seq	F	M00054802:72	CH17COHLV
284	452567	RTA22200002F.j.12.1.P.Seq	F	M00055519:36	CH15CON
285	411113	RTA22200024F.f.03.1.P.Seq	F	M00055185:21	CH17COHLV
286	650749	RTA22200002F.d.18.1.P.Seq	F	M00055451:611	CH15CON
287	558899	RTA22200026F.d.12.1.P.Seq	F	M00055421:44	CH17COHLV
288	452986	RTA22200002F.a.13.1.P.Seq	F	M00055426:22	CH15CON
289	393197	RTA22200015F.k.01.1.P.Seq	F	M00057108:59	CH16COP
290	499424	RTA22200024F.e.10.1.P.Seq	F	M00055179:42	CH17COHLV
291	21669	RTA22200025F.m.07.2.P.Seq	F	M00055175:42	CH17COHLV
292	640590	RTA22200004F.h.21.1.P.Seq	F	M00055794:711	CH15CON
293	549936	RTA22200004F.c.10.1.P.Seq	F	M00055754:711	
294	448770	RTA222000241.c.10.1.1.1.Seq	F	M00053137:311	CH16COP
295	559280	RTA22200015F.h.14.1.P.Seq	F	M00057174.712	CH16COP
296	648934	RTA22200013F.n.14.1.F.Seq	F	M00057093.09 M00055591:81	CH15CON
297	452685	RTA22200003F.C.10.1.F.Seq	F	M00055838:412	CH15CON
298	456549	RTA22200004F.p.00.1.F.Seq	F	M00055522:32	CH15CON
290 299	446614	RTA22200002F.j.24.1.P.Seq	F	M00033322.32 M00042563:52	CH15CON
300	559280	RTA22200001F.d.24.1.F.Seq	F	M00042303.52 M00055170:52	CH17COHLV
300	446673	RTA22200024F.d.16.1.F.Seq	F	M00057055:78	CH16COP
302	562550	RTA22200013F.C.01.1.F.Seq	F	M00057055.78	CH15CON
303	467288	RTA22200004F.II.17.1.F.Seq	F	M00055500:25	CH15CON
303	467288	RTA22200002F.II.10.1.P.Seq	F		
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	······································	F	M00055823:43	CH15CON
305 306	393197 407077	RTA22200015F.j.24.1.P.Seq	F	M00057108:59	CH16COP CH16COP
		RTA22200015F.j.22.1.P.Seq	F F	M00057108:64	Annual Communication of the co
307 308	499424	RTA22200018F.h.13.1.P.Seq RTA22200021F.o.20.2.P.Seq	F F	M00043317:81	CH17COHLV
~*~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	554500			M00054857:512	
309	730143	RTA22200015F.k.14.1.P.Seq	F F	M00057112:29	CH16COP
310	595506	RTA22200015F.c.21.1.P.Seq	F	M00057061:44	CH16COP
311	2334	RTA22200016F.k.13.1.P.Seq	vanamente automoranismo antigente an	M00057242:85	CH16COP
312	647444	RTA22200002F.k.22.1.P.Seq	F	M00055527:711	CH15CON
313	380291	RTA22200008F.a.03.1.P.Seq	E	M00056292:55	CH15CON
314	644849	RTA22200016F.a.22.1.P.Seq	transcription of the contract	M00057162:37	CH16COP
315	449457	RTA22200018F.g.23.1.P.Seq	<u> </u>	M00043314:84	CH17COHLV
316	446673	RTA22200015F.b.24.1.P.Seq	<u> </u>	M00057055:78	CH16COP
317	549069	RTA22200020F.e.05.1.P.Seq	<u>F</u>	M00054596:27	CH17COHLV
318	728884	RTA22200015F.d.02.1.P.Seq	<u> </u>	M00057063:38	CH16COP
319	415058	RTA22200020F.f.03.1.P.Seq		M00054605:41	CH17COHLV
320	553955	RTA22200020F.e.20.1.P.Seq	F_	M00054602:42	CH17COHLV
321	455820	RTA22200018F.i.16.1.P.Seq	<u> </u>	M00043323:44	CH17COHLV
322	549617	RTA22200020F.f.20.1.P.Seq	<u>F</u>	M00054611:62	CH17COHLV
323	449831	RTA22200018F.d.22.1.P.Seq	F	M00042518:16	CH17COHLV
324	451580	RTA22200018F.b.09.1.P.Seq	F	M00042444:75	CH17COHLV

Table 1

r	ble 1		,		
SEQ ID	MORTHUM GOL 1 - PERSON 14 probabilitations	SEQ NAME	ORIENTATION	and a superior of the second s	LIBRARY
325	558899	RTA22200025F.f.09.1.P.Seq	F	M00055325:212	CH17COHLV
326	562292	RTA22200025F.j.10.1.P.Seq	F	M00055368:26	CH17COHLV
327	5830	RTA22200011F.d.11.1.P.Seq	F	M00056552:110	CH16COP
328	8953	RTA22200248F.p.09.1.P.Seq	F	M00027608:87	CH04MAL
329	8012	RTA22200233F.i.02.1.P.Seq	F	M00008089:59	CH03MAH
330	185718	RTA22200250F.f.15.1.P.Seq	F	M00027829:42	CH04MAL
331	729851	RTA22200011F.b.18.1.P.Seq	F	M00056537:85	CH16COP
332	185597	RTA22200249F.a.20.1.P.Seq	F	M00027616:712	CH04MAL
333	9887	RTA22200234F.h.14.1.P.Seq	F	M00022253:53	CH03MAH
334	725825	RTA22200011F.c.16.1.P.Seq	F	M00056547:34	CH16COP
335	6545	RTA22200249F.e.12.1.P.Seq	F	M00027639:511	CH04MAL
336	21205	RTA22200249F.f.24.1.P.Seq	F	M00027660:53	CH04MAL
337	8867	RTA22200234F.j.23.1.P.Seq	F	M00022280:711	CH03MAH
338	729295	RTA22200010F.m.09.1.P.Seq	F	M00056499:65	CH16COP
339	730430	RTA22200010F.h.08.1.P.Seq	F	M00056416:212	CH16COP
340	7072	RTA22200233F.h.20.1.P.Seq	F ;	M00008085:39	CH03MAH
341	730533	RTA22200010F.n.17.1.P.Seq	F	M00056506:712	CH16COP
342	9121	RTA22200224F.g.11.1.P.Seq	F	M00005312:410	CH02COH
343	11131	RTA22200224F.c.08.1.P.Seq	F	M00004852:14	CH02COH
344	640116	RTA22200010F.n.01.1.P.Seq	F	M00056503:711	CH16COP
345	730282	RTA22200010F.k.18.1.P.Seq	F	M00056483:66	CH16COP
346	550571	RTA22200019F.h.11.1.P.Seq	F	M00054502:52	CH17COHLV
347	1183	RTA22200248F.c.22.1.P.Seq	F	M00027527:74	CH04MAL
348	449437	RTA22200019F.g.09.1.P.Seq	F	M00054494:51	CH17COHLV
349	8966	RTA22200242F.d.21.1.P.Seq	F	M00026994:17	CH04MAL
350	6134	RTA22200006F.p.04.2.P.Seq	F	M00056125:49	CH15CON
351	95700	RTA22200241F.f.09.1.P.Seq	F	M00026896:510	CH04MAL
352	7066	RTA22200229F.a.23.1.P.Seq	F	M00006928:14	CH02COH
353	648310	RTA22200007F.a.23.1.P.Seq	F	M00056143:59	CH15CON
354	730059	RTA22200013F.k.11.1.P.Seq	F	M00056874:45	CH16COP
355	736014	RTA22200012F.f.11.1.P.Seq	F ;	M00056708:411	CH16COP
356	646577	RTA22200012F.d.07.1.P.Seq	F	M00056693:38	CH16COP
357	732254	RTA22200014F.g.09.1.P.Seq	F :	M00056967:57	CH16COP
358	7037	RTA22200229F.b.01.1.P.Seq	,	M00006928:41	CH02COH
359	7037	RTA22200229F.a.24.1.P.Seq	F	M00006928:41	CH02COH
360	6937	RTA22200232F.b.06.1.P.Seq	F	M00021864:57	CH03MAH
361	7572	RTA22200228F.k.06.2.P.Seq		M00006867:612	CH02COH
362	388085	RTA22200005F.c.21.1.P.Seq		M00055868:43	CH15CON
363	2676	RTA22200227F.p.19.1.P.Seq	F	M00006761:35	CH02COH
364	639240	RTA22200007F.k.18.1.P.Seq	F	M00056238:57	CH15CON
365	650472	RTA22200012F.e.19.1.P.Seq	F	M00056705:57	CH16COP
366	727789	RTA22200012F.g.19.1.P.Seq	F	M00056715:31	CH16COP
367	2495	RTA22200238F.e.03.1.P.Seq	F	M00022895:211	CH03MAH
368	732254	RTA22200014F.g.09.2.P.Seq	F	M00056967:57	CH16COP
369	5268	RTA22200225F.m.13.1.P.Seq	F	M00005517:64	CH02COH
370	11881	RTA22200238F.e.09.1.P.Seq	F	M00022897:83	CH03MAH
371	448677	RTA22200009F.b.05.2.P.Seq	F ,	M00042440:55	CH16COP
372	1876	RTA222000091.b.03.2.17.Seq	F	M00042446:64	CH03MAH
373	3441	RTA22200233F.e.23.1.P.Seq	F	M000022490.04 M00008065:62	CH03MAH
374	726134	RTA22200233F.E.23.1.F.Seq	**************************************	M00056618:611	CH16COP
375	9048	RTA22200011F.1.03.1.F.Seq	F	M00003804:19	CH01COH
376	26489	RTA22200242F.k.07.1.P.Seq		M00003804.19	CH04MAL
377	644205	RTA22200007F.c.24.1.P.Seq	F	M00027039.88	CH15CON
378	468689	RTA22200007F.C.24.1.F.Seq		M00056036:26	CH15CON CH15CON
010	+00000	17174420000007.1.04.2.7.389	F ,	19100000000.20	VILIOCON

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	ble 1				
	TO THE RESIDENCE AND ADDRESS ASSESSMENT OF THE PARTY OF T	SEQ NAME	ORIENTATION	CARLO CONTRACTOR SECURIO CONTRACTOR DE LA CONTRACTOR DE L	LIBRARY
379	638971	RTA22200012F.k.19.1.P.Seq	F	M00056746:12	CH16COP
380	10274	RTA22200241F.I.17.1.P.Seq	F	M00026936:312	CH04MAL
381	6725	RTA22200232F.f.22.1.P.Seq	F	M00022013:85	CH03MAH
382	2488	RTA22200012F.I.21.1.P.Seq	F	M00056754:49	CH16COP
383	8366	RTA22200244F.c.08.1.P.Seq	F	M00027173:48	CH04MAL
384	502683	RTA22200007F.a.09.1.P.Seq	F	M00056139:71	CH15CON
385	450914	RTA22200012F.k.22.1.P.Seq	F	M00056747:45	CH16COP
386	21205	RTA22200243F.k.21.1.P.Seq	F	M00027140:311	CH04MAL
387	644205	RTA22200007F.d.01.1.P.Seq	F	M00056162:68	CH15CON
388	5268	RTA22200225F.m.05.1.P.Seq	F	M00005513:69	CH02COH
389	8012	RTA22200232F.n.17.1.P.Seq	F	M00022148:16	CH03MAH
390	11270	RTA22200227F.m.18.1.P.Seq	F	M00006734:18	CH02COH
391	10924	RTA22200237F.h.04.1.P.Seq	F	M00022738:46	CH03MAH
392	11619	RTA22200241F.g.14.1.P.Seq	F	M00026902:74	CH04MAL
393	3650	RTA22200236F.e.11.1.P.Seq	F	M00022617:32	CH03MAH
394	1655	RTA22200222F.e.14.1.P.Seq	F	M00001637:49	CH01COH
395	3275	RTA22200238F.b.21.1.P.Seq	F	M00022876:25	CH03MAH
396	3355	RTA22200238F.c.05.1.P.Seq	F	M00022880:79	CH03MAH
397	2078	RTA22200235F.g.18.1.P.Seq	F	M00022473:26	СНОЗМАН
398	4809	RTA22200222F.g.09.1.P.Seq	F	M00003794:47	CH01COH
399	6402	RTA22200236F.I.15.1.P.Seq	F	M00022660:24	CH03MAH
400	555244	RTA22200023F.e.03.1.P.Seq	F	M00055042:21	CH17COHLV
401	548965	RTA22200012F.g.16.1.P.Seq	F	M00056715:410	CH16COP
402	4747	RTA22200227F.m.12.1.P.Seq	F	M00006731:38	CH02COH
403	40208	RTA22200241F.n.21.1.P.Seq	F	M00026950:81	CH04MAL
404	14596	RTA22200241F.d.18.1.P.Seq	F	M00026879:22	CH04MAL
405	7110	RTA22200232F.n.01.1.P.Seq	F	M00022143:41	CH03MAH
406	7110	RTA22200232F.m.24.1.P.Seq	F	M00022143:41	CH03MAH
407	6592	RTA22200238F.p.20.1.P.Seq	F	M00023029:56	CH03MAH
408	6455	RTA22200232F.o.06.1.P.Seq	F	M00022151:75	CH03MAH
409	2738	RTA22200232F.g.16.1.P.Seq	F	M00022050:44	CH03MAH
410	696	RTA22200232F.m.02.1.P.Seq	F	M00022132:64	СНОЗМАН
411	379186	RTA22200012F.c.05.1.P.Seq	F	M00056682:610	CH16COP
412	1588	RTA22200232F.o.03.1.P.Seq	F	M00022151:411	CH03MAH
413	7007	RTA22200225F.e.05.1.P.Seq	F	M00005454:33	CH02COH
414	9025	RTA22200222F.k.23.1.P.Seq	F	M00004080:15	CH01COH
415	650749	RTA22200007F.i.16.2.P.Seq	F	M00056220:49	CH15CON
416	553158	RTA22200005F.e.13.1.P.Seq	F	M00055879:511	CH15CON
417	641703	RTA22200003F.e.18.1.P.Seq	F	M00055628:18	CH15CON
418	833	RTA22200006F.j.24.2.P.Seq	F	M00056077:512	CH15CON
419	649259	RTA22200006F.o.01.2.P.Seq	F	M00056108:212	CH15CON
420	451179	RTA22200011F.i.14.1.P.Seq	F	M00056596:88	CH16COP
421	9505	RTA22200231F.b.20.1.P.Seq	F I	M00007935:15	CH03MAH
422	736728	RTA22200015F.n.19.1.P.Seq	F	M00057134:31	CH16COP
423	380412	RTA22200001F.p.23.1.P.Seq	F	M00054918:43	CH15CON
424	642425	RTA22200011F.k.12.1.P.Seq	F	M00056613:15	CH16COP
425	405073	RTA2220007F.o.05.1.P.Seq	F	M00056273:11	CH15CON
426	174250	RTA222000071.0.00.111.0cq	F	M00056475:61	CH15CON
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428	639029	RTA222000171.d.00.1.11.Seq	F	M00057524.412	CH15CON
429	452245	RTA222000071.R.05.2.F.Seq	F	M00054963:811	CH17COHLV
430	510254	RTA22200022F.i.03.1.F.Seq	F	M00054983.811 M00055992:511	CH15CON
431	642425	RTA22200003F.p.22.1.F.3eq	F	M00055992.511	CH15CON
432	51939	RTA22200007F.g.19.1.F.Seq	F	M00055912:510	CH15CON
	J 1 3 J 3	1172220000011.1.22.1.F.389	I i	141000003 12.010	CITIOCON

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	DIE I	OFO NAME	ODIENTATION	OLONE ID	LIDEADY
contraction and the property and the party of	CLUSTER	SEQ NAME	ORIENTATION	THE PARTY OF THE P	LIBRARY
433	7379	RTA22200228F.f.21.1.P.Seq		M00006821:21	CH02COH
434	546632	RTA22200013F.h.16.1.P.Seq	Ę	M00056858:112	CH16COP
435	734827	RTA22200011F.m.09.1.P.Seq	<u> </u>	M00056631:38	CH16COP
436	2554	RTA22200231F.g.06.1.P.Seq	F_	M00007961:65	CH03MAH
437	643285	RTA22200012F.a.16.1.P.Seq	F	M00056665:111	CH16COP
438	448770	RTA22200001F.o.07.1.P.Seq	F	M00054793:27	CH15CON
439	375380	RTA22200023F.e.20.1.P.Seq	<u> </u>	M00055046:37	CH17COHLV
440	726134	RTA22200011F.l.10,1.P.Seq	F	M00056620:512	CH16COP
441	422687	RTA22200015F.p.03.1.P.Seq	<b>F</b>	M00057143:55	CH16COP
442	448436	RTA22200012F.n.02.1.P.Seq	F	M00056763:45	CH16COP
443	644893	RTA22200012F.I.24.1.P.Seq	F	M00056754:15	CH16COP
444	559104	RTA22200012F.n.21.1.P.Seq	F	M00056771:312	CH16COP
445	551172	RTA22200016F.g.09.1.P.Seq	F	M00057211:16	CH16COP
446	724296	RTA22200012F.f.22.1.P.Seq	F	M00056710:67	CH16COP
447	735936	RTA22200009F.p.13.1.P.Seq	F	M00056346:312	CH16COP
448	556326	RTA22200023F.k.20.1.P.Seq	F	M00055085:110	CH17COHLV
449	729699	RTA22200011F.o.04.1.P.Seq	F	M00056646:32	CH16COP
450	550694	RTA22200022F.a.05.1.P.Seq	F	M00054867:27	CH17COHLV
451	734738	RTA22200017F.e.11.1.P.Seq	F	M00057337:72	CH16COP
452	404502	RTA22200007F.p.08.1.P.Seq	F	M00056286:58	CH15CON
453	554151	RTA22200015F.o.18.1.P.Seq	F	M00057142:17	CH16COP
454	649852	RTA22200001F.n.24.1.P.Seq	F	M00042914:210	CH15CON
455	734063	RTA22200011F.m.16.1.P.Seq	F	M00056633:27	CH16COP
456	7279	RTA22200230F.b.22.1.P.Seq	F	M00007108:41	CH02COH
457	2676	RTA22200230F.I.05.1.P.Seq	F	M00007204:41	CH02COH
458	649148	RTA22200006F.k.08.2.P.Seq	F	M00056079:311	CH15CON
459	1953	RTA22200226F.p.21.1.P.Seq	F	M00006576:24	CH02COH
460	650108	RTA22200006F.d.04.2.P.Seq	F	M00056018:75	CH15CON
461	515350	RTA22200005F.g.04.1.P.Seq	F	M00055886:79	CH15CON
462	402494	RTA22200013F.a.15.1.P.Seq	F	M00056796:85	CH16COP
463	649148	RTA22200013F.d.09.1.P.Seq	F	M00056821:39	CH16COP
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466	453079	RTA22200009F.k.11.2.P.Seq	F	M00042835:42	CH16COP
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468	644903	RTA22200003F.f.09.1.P.Seq	F	M00055635:74	CH15CON
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Table 1

	ble 1		<del>,                                    </del>		
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490	612572	RTA22200007F.I.08.2.P.Seq	F	M00056244:28	CH15CON
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Samuel and the second	158601	RTA22200222F.k.16.1.F.Seq	<u> </u>	M00004000.42	CH04MAL
532	By and a second contract of the contract of th		F		CH04MAL
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Table 1

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588	406413	RTA22200021F.c.09.2.P.Seq	F	M00054734:810	<u>จุ้าวางสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถ</u>
589	555103	RTA22200024F.a.21.1.P.Seq	F	M00055148:411	CH17COHLV
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592	727181	RTA22200016F.k.12.1.P.Seq	F	M00057242:29	CH16COP
593	551117	RTA22200021F.f.14.3.P.Seq	F	M00054760:112	Santonia mariante de como con como con como con como como co
594	464040	RTA22200024F.g.21.1.P.Seq	F	M00055198:77	CH17COHLV

Table 1

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600	549722	RTA22200020F.a.09.1.P.Seq	<u>F</u>	M00054568:711	
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606	553802	RTA22200022F.f.04.1.P.Seq	F	M00054908:67	CH17COHLV
607	639662	RTA22200004F.h.15.1.P.Seq	F	M00055792:79	CH15CON
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612	417155	RTA22200002F.n.24.1.P.Seq	F	M00055548:512	CH15CON
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625	451092	RTA22200026F.h.02.1.P.Seq	F	M00055487:66	CH17COHLV
626	546642	RTA22200016F.o.16.1.P.Seq	F	M00057275:112	CH16COP
627	553736	RTA22200022F.j.18.1.P.Seq	F	M00054952:61	CH17COHLV
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Table 1

A94   A91441   RTA22200003F b.12.1 P.Seq	SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
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652   551907   RTA22200020F.I.04.1.P.Seq   F   M0005578-3.73   CH15CONLO   654   447532   RTA22200003F.b.14.1.P.Seq   F   M0005578-3.73   CH15CON   655   558454   RTA22200026F.c.17.1.P.Seq   F   M0005584-311   CH15CON   656   502883   RTA22200026F.c.17.1.P.Seq   F   M0005487-82   CH17COHLV   657   446909   RTA22200023F.n.17.1.P.Seq   F   M0005487-82   CH17COHLV   658   452506   RTA22200023F.c.02.1.P.Seq   F   M0005487-89   CH17COHLV   659   449792   RTA22200023F.c.02.1.P.Seq   F   M0005487-89   CH17COHLV   650   549395   RTA22200020F.i.08.1.P.Seq   F   M00055027-88   CH17COHLV   651   243653   RTA22200020F.i.08.1.P.Seq   F   M00054838-49   CH17COHLV   662   445911   RTA22200021F.s.01.P.Seq   F   M00054838-49   CH17COHLV   663   445031   RTA22200001F.s.18.1.P.Seq   F   M00054821-36   CH17COHLV   664   431032   RTA22200018F.g.04.1.P.Seq   F   M00043251-36   CH17COHLV   665   446860   RTA22200018F.g.04.1.P.Seq   F   M00043310-23   CH17COHLV   666   446860   RTA22200018F.g.04.1.P.Seq   F   M00043310-23   CH17COHLV   666   641884   RTA22200018F.g.04.1.P.Seq   F   M00043310-33   CH17COHLV   667   452800   RTA22200018F.g.04.1.P.Seq   F   M00043310-36   CH17COHLV   668   641884   RTA222000018F.g.04.1.P.Seq   F   M00043301-36   CH17COHLV   669   548965   RTA222000018F.g.04.1.P.Seq   F   M00043301-36   CH17COHLV   669   669   RTA22000018F.g.04.1.P.Seq   F   M0004300-30-324   CH15CON   667   452800   RTA222000018F.g.08.1.P.Seq   F   M0004300-30-324   CH15CON   667   452800   RTA222000018F.g.08.1.P.Seq   F   M0004571-71-81   CH17COHLV   669   669   RTA220000018F.g.08.1.P.Seq   F   M00056771-816   CH17COHLV   670   734793   RTA222000018F.g.08.1.P.Seq   F   M00056774-81   CH15CON   671   539955   RTA222000020F.s.08.1.P.Seq   F   M00054596-71   CH17COHLV   672   561992   RTA2220000218F.g.08.1.P.Seq   F   M00054596-71   CH17COHLV   673   562292   RTA2220000218F.g.08.1.P.Seq   F   M00054596-71   CH17COHLV   674   420886   RTA222000018F.g.08.1.P.Seq   F   M00054596-71   CH17COHLV   675   64346   RTA222000018F.g.08.1.P.Seq   F	<u> </u>	&				Emmanue de la companya del la companya de la compan
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655         558454         RTA22200026F.c.17.1.P.Seq         F         M00056447:82         CH17COHLV           657         446909         RTA22200029F.j.12.1.P.Seq         F         M00055112:33         CH17COHLV           658         452506         RTA22200029F.c.23.1.P.Seq         F         M0005512:83         CH17COHLV           659         449792         RTA22200029F.c.03.1.P.Seq         F         M00055027:88         CH17COHLV           660         549395         RTA22200029F.io.8.1.P.Seq         F         M00055627:36         CH17COHLV           661         234653         RTA22200029F.jo.8.1.P.Seq         F         M00054621:36         CH17COHLV           662         453911         RTA22200018F.jo.1.P.Seq         F         M00045621:36         CH17COHLV           663         452071         RTA22200018F.jo.4.1.P.Seq         F         M0004331:36         CH17COHLV           664         451032         RTA22200018F.jo.8.1.P.Seq         F         M0004331:36         CH17COHLV           665         44680         RTA22200008F.jo.8.1.P.Seq         F         M0004331:36         CH17COHLV           666         44184         RTA22200004F.jo.8.1.P.Seq         F         M0004331:3:36         CH17COHLV           667	-	gano moran marana mang	Augustus Aug		THE PROPERTY OF THE PROPERTY AND ADDRESS OF THE PROPERTY OF TH	Assertance of the contract of
655   502883   RTA22200020F_i.12.1.P.Seq   F   M00054647:82 CH17COHLV   657   446909   RTA22200023F.n.17.1.P.Seq   F   M00055112:33 CH17COHLV   658   452506   RTA22200023F.c.23.1.P.Seq   F   M00055457:69 CH17COHLV   659   449792   RTA22200023F.c.23.1.P.Seq   F   M00055627:68 CH17COHLV   660   549395   RTA22200020F.i.08.1.P.Seq   F   M0005663*49 CH17COHLV   661   234653   RTA22200020F.g.14.1.P.Seq   F   M00054621:36 CH17COHLV   662   453911   RTA22200015F.f.01.1.P.Seq   F   M00054621:36 CH17COHLV   663   452071   RTA22200015F.f.01.1.P.Seq   F   M00045252*77 CH15CON   664   451032   RTA22200018F.g.04.1.P.Seq   F   M00043310:23 CH17COHLV   665   446860   RTA22200018F.g.17.1.P.Seq   F   M00043310:23 CH17COHLV   666   446860   RTA22200018F.g.17.1.P.Seq   F   M00043310:33 CH17COHLV   667   452800   RTA22200018F.b.08.1.P.Seq   F   M00043310:36 CH17COHLV   668   4481835   RTA22200004F.i.08.1.P.Seq   F   M000560503:24 CH15CON   669   548965   RTA22200004F.i.08.1.P.Seq   F   M0005560778:69 CH15CON   670   734793   RTA222000026F.g.08.1.P.Seq   F   M000556778:69 CH15CON   671   539955   RTA222000026F.g.12.1.P.Seq   F   M00054609:61 CH17COHLV   673   556292   RTA22200020F.s.12.1.P.Seq   F   M00054609:61 CH17COHLV   674   420866   RTA22200004F.i.p.Seq   F   M00054699:61 CH17COHLV   675   9436   RTA22200004F.i.p.Seq   F   M00054699:61 CH17COHLV   676   44424   RTA22200004F.i.p.Seq   F   M00054699:61 CH17COHLV   677   412364   RTA22200004F.i.p.Seq   F   M00054596:711 CH17COHLV   678   44424   RTA22200004F.i.p.Seq   F   M00055074:31 CH15CON   679   9436   RTA22200004F.i.p.Seq   F   M00055074:31 CH15CON   679   679   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670   670	\$	şş	un aproximation and a proximation and a proximation of the contract of the con			\$~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
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658         452566         RTA22200020F.c.02.1.P.Seq         F         M000556378.69         CH17COHLV           669         449792         RTA22200021F.c.02.1.P.Seq         F         M0005638.49         CH17COHLV           660         549395         RTA22200020F.g.14.1.P.Seq         F         M00054638.49         CH17COHLV           661         234653         RTA22200001F.g.14.1.P.Seq         F         M00054621.36         CH17COHLV           662         453911         RTA2220001F.g.14.1.P.Seq         F         M00045259:77         CH15CON           663         452071         RTA2220001F.g.04.1.P.Seq         F         M0004320:23         CH15CON           664         4451032         RTA2220001F.g.04.1.P.Seq         F         M00043310:36         CH15CON           665         44680         RTA22200001F.g.16.1.P.Seq         F         M00056303:24         CH15CON           666         641884         RTA22200001F.g.08.1.P.Seq         F         M00056303:24         CH15CON           667         452800         RTA22200001F.c.08.1.P.Seq         F         M0005605778:69         CH15CON           669         548965         RTA22200002F.e.08.1.P.Seq         F         M00056056771-19         CH15CON           671         53	3	&				ån
659         449792         RTA22200029F.i.08.1.P.Seq         F         M00055027-68         CH17COHLV           660         549395         RTA22200020F.i.08.1.P.Seq         F         M00054681-36         CH17COHLV           661         234653         RTA22200020F.g.08.1.P.Seq         F         M00057074-39         CH16COPL           662         453911         RTA22200018F.g.01.1.P.Seq         F         M00047074-39         CH16COPL           663         452071         RTA22200018F.g.01.1.P.Seq         F         M00043310:33         CH17COHLV           664         451032         RTA22200018F.g.17.1.P.Seq         F         M00043310:33         CH17COHLV           665         44680         RTA22200018F.g.17.1.P.Seq         F         M00043310:33         CH17COHLV           666         641884         RTA22200008F.g.08.1.P.Seq         F         M00043301:66         CH15CON           667         452800         RTA22200020F.g.08.1.P.Seq         F         M000540778:89         CH15CON           669         548965         RTA22200020F.g.08.1.P.Seq         F         M0005407174:812         CH16CON           671         539955         RTA22200009F.g.06.1.P.Seq         F         M0005409:41         CH16CON           672         <	Annual Company of the	American and a superior and a superi	and the second s	·	Contraction of the contraction	Emmironementen semeraturatur armeniaturatur elementen el
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668         461835         RTA22200020F.g.08.1.P.Seq         F         M00055778:69         CH15CON           669         548965         RTA22200020F.g.08.1.P.Seq         F         M00054617:19         CH17COHLV           670         734793         RTA22200016F.c.20.1.P.Seq         F         M00057174:812         CH16COP           671         539955         RTA22200020F.c.06.1.P.Seq         F         M0005469:61         CH15CON           672         561892         RTA22200020F.f.11.1.P.Seq         F         M00054596:71         CH17COHLV           673         562292         RTA22200001F.j.02.1.P.Seq         F         M0005578:35         CH15CON           674         420686         RTA22200004F.j.20.1.P.Seq         F         M0005578:35         CH15CON           675         9436         RTA22200004F.j.20.1.P.Seq         F         M00055774:51         CH15CON           676         1013         RTA22200004F.j.20.1.P.Seq         F         M00055373:33         CH15CON           677         412364         RTA22200004F.j.j.18.1.P.Seq         F         M00043327:82         CH17COHLV           679         394413         RTA22200015F.j.1.P.Seq         F         M00043347:712         CH17COHLV           679         394413 <td>Secretaria de la companya del companya del companya de la companya</td> <td>&amp;</td> <td></td> <td></td> <td></td> <td>Samera a construence de la construence del la construence del la construence de la c</td>	Secretaria de la companya del companya del companya de la companya	&				Samera a construence de la construence del la construence del la construence de la c
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681         455032         RTA22200018F.I.07.1.P.Seq         F         M00043347:712         CH17COHLV           682         185400         RTA22200025F.b.17.1.P.Seq         F         M00055279:512         CH17COHLV           683         453911         RTA22200015F.e.24.1.P.Seq         F         M00057074:39         CH16COP           684         650297         RTA22200004F.e.23.1.P.Seq         F         M00055774:73         CH15CON           685         185400         RTA22200018F.k.02.1.P.Seq         F         M00043338:23         CH17COHLV           686         449512         RTA22200025F.c.06.1.P.Seq         F         M00055283:82         CH17COHLV           687         44424         RTA22200018F.i.24.1.P.Seq         F         M00043327:82         CH17COHLV           688         556216         RTA22200025F.h.09.1.P.Seq         F         M00055324:810         CH17COHLV           689         448677         RTA22200025F.e.16.1.P.Seq         F         M00055319:11         CH17COHLV           690         375380         RTA22200025F.b.03.1.P.Seq         F         M00055319:11         CH17COHLV           691         376988         RTA22200025F.g.20.1.P.Seq         F         M00055346:42         CH17COHLV           693	<u> </u>	Emmercania and a second			*************************************	
682         185400         RTA22200025F.b.17.1.P.Seq         F         M00055279:512         CH17COHLV           683         453911         RTA22200015F.e.24.1.P.Seq         F         M00057074:39         CH16COP           684         650297         RTA2220004F.e.23.1.P.Seq         F         M00055774:73         CH15CON           685         185400         RTA22200018F.k.02.1.P.Seq         F         M00043338:23         CH17COHLV           686         449512         RTA22200025F.c.06.1.P.Seq         F         M00043327:82         CH17COHLV           687         44424         RTA22200018F.i.24.1.P.Seq         F         M00043327:82         CH17COHLV           688         556216         RTA22200025F.h.09.1.P.Seq         F         M00055330:61         CH17COHLV           689         448677         RTA22200025F.e.16.1.P.Seq         F         M00055319:11         CH17COHLV           690         375380         RTA22200025F.e.16.1.P.Seq         F         M00055319:11         CH17COHLV           691         379341         RTA22200025F.e.920.1.P.Seq         F         M00055344:81         CH17COHLV           692         376988         RTA22200025F.g.20.1.P.Seq         F         M00055344:89         CH17COHLV           693	<u> </u>	\$			~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
683         453911         RTA22200015F.e.24.1.P.Seq         F         M00057074:39         CH16COP           684         650297         RTA22200004F.e.23.1.P.Seq         F         M00055774:73         CH15CON           685         185400         RTA22200018F.k.02.1.P.Seq         F         M00043338:23         CH17COHLV           686         449512         RTA22200025F.c.06.1.P.Seq         F         M00055283:82         CH17COHLV           687         44424         RTA22200018F.i.24.1.P.Seq         F         M00043327:82         CH17COHLV           688         556216         RTA22200025F.h.09.1.P.Seq         F         M00055350:61         CH17COHLV           689         448677         RTA22200025F.f.07.1.P.Seq         F         M00055324:810         CH17COHLV           690         375380         RTA22200025F.e.16.1.P.Seq         F         M00055319:11         CH17COHLV           691         379341         RTA22200018F.a.24.1.P.Seq         F         M00055346:42         CH17COHLV           692         376988         RTA22200025F.b.03.1.P.Seq         F         M00055346:42         CH17COHLV           693         559806         RTA22200025F.g.20.1.P.Seq         F         M00055368:310         CH17COHLV           695	-	-			**************************************	
684         650297         RTA22200004F.e.23.1.P.Seq         F         M00055774:73         CH15CON           685         185400         RTA22200018F.k.02.1.P.Seq         F         M00043338:23         CH17COHLV           686         449512         RTA22200025F.c.06.1.P.Seq         F         M00055283:82         CH17COHLV           687         44424         RTA22200018F.i.24.1.P.Seq         F         M00043327:82         CH17COHLV           688         556216         RTA22200025F.h.09.1.P.Seq         F         M00055350:61         CH17COHLV           689         448677         RTA22200025F.e.16.1.P.Seq         F         M00055319:11         CH17COHLV           690         375380         RTA22200025F.e.16.1.P.Seq         F         M00055319:11         CH17COHLV           691         379341         RTA22200018F.a.24.1.P.Seq         F         M00055346:42         CH17COHLV           692         376988         RTA22200025F.b.03.1.P.Seq         F         M00055344:89         CH17COHLV           693         559806         RTA22200025F.j.07.1.P.Seq         F         M00055348:89         CH17COHLV           694         550195         RTA22200025F.k.09.1.P.Seq         F         M00055348:89         CH17COHLV           695	\$	&		3		Committee and the committee of the commi
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688         556216         RTA22200025F.h.09.1.P.Seq         F         M00055350:61         CH17COHLV           689         448677         RTA22200025F.f.07.1.P.Seq         F         M00055324:810         CH17COHLV           690         375380         RTA22200025F.e.16.1.P.Seq         F         M00055319:11         CH17COHLV           691         379341         RTA22200018F.a.24.1.P.Seq         F         M00042442:12         CH17COHLV           692         376988         RTA22200025F.h.03.1.P.Seq         F         M00055346:42         CH17COHLV           693         559806         RTA22200025F.g.20.1.P.Seq         F         M00055344:89         CH17COHLV           694         550195         RTA22200025F.k.09.1.P.Seq         F         M00055368:310         CH17COHLV           695         562221         RTA22200025F.k.09.1.P.Seq         F         M00055374:18         CH17COHLV           696         211         RTA22200248F.o.19.1.P.Seq         F         M00027607:85         CH04MAL           697         6751         RTA22200248F.o.12.1.P.Seq         F         M00027604:710         CH04MAL           699         11619         RTA22200249F.d.17.1.P.Seq         F         M00027608:5         CH04MAL           701 <td< td=""><td>§</td><td></td><td>туруны тоо торуу жана тоо тоо уурын жана байган айын байган байг</td><td></td><td>**************************************</td><td></td></td<>	§		туруны тоо торуу жана тоо тоо уурын жана байган айын байган байг		**************************************	
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691         379341         RTA22200018F.a.24.1.P.Seq         F         M00042442:12         CH17COHLV           692         376988         RTA22200025F.h.03.1.P.Seq         F         M00055346:42         CH17COHLV           693         559806         RTA22200025F.g.20.1.P.Seq         F         M00055344:89         CH17COHLV           694         550195         RTA22200025F.j.07.1.P.Seq         F         M00055368:310         CH17COHLV           695         562221         RTA22200025F.k.09.1.P.Seq         F         M00055374:18         CH17COHLV           696         211         RTA22200248F.o.19.1.P.Seq         F         M00027607:85         CH04MAL           697         6751         RTA22200248F.o.01.1.P.Seq         F         M00027604:710         CH04MAL           698         6751         RTA22200248F.n.24.1.P.Seq         F         M00027604:710         CH04MAL           699         11619         RTA22200250F.d.17.1.P.Seq         F         M00027717:75         CH04MAL           700         2883         RTA22200249F.i.14.1.P.Seq         F         M00027628:11         CH04MAL           701         9784         RTA22200249F.c.12.1.P.Seq         F         M00027628:11         CH04MAL	Sergeneering	and the second s			GERT MANNES AND A TONOMORE STATE OF THE AND A STATE	
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693         559806         RTA22200025F.g.20.1.P.Seq         F         M00055344:89         CH17COHLV           694         550195         RTA22200025F.j.07.1.P.Seq         F         M00055368:310         CH17COHLV           695         562221         RTA22200025F.k.09.1.P.Seq         F         M00055374:18         CH17COHLV           696         211         RTA22200248F.o.19.1.P.Seq         F         M00027607:85         CH04MAL           697         6751         RTA22200248F.o.01.1.P.Seq         F         M00027604:710         CH04MAL           698         6751         RTA22200248F.n.24.1.P.Seq         F         M00027604:710         CH04MAL           699         11619         RTA22200250F.d.17.1.P.Seq         F         M00027806:85         CH04MAL           700         2883         RTA22200249F.I.14.1.P.Seq         F         M00027717:75         CH04MAL           701         9784         RTA22200249F.c.12.1.P.Seq         F         M00027628:11         CH04MAL	Section of the sectio				·	
694         550195         RTA22200025F.j.07.1.P.Seq         F         M00055368:310         CH17COHLV           695         562221         RTA22200025F.k.09.1.P.Seq         F         M00055374:18         CH17COHLV           696         211         RTA22200248F.o.19.1.P.Seq         F         M00027607:85         CH04MAL           697         6751         RTA22200248F.o.01.1.P.Seq         F         M00027604:710         CH04MAL           698         6751         RTA22200248F.n.24.1.P.Seq         F         M00027604:710         CH04MAL           699         11619         RTA22200250F.d.17.1.P.Seq         F         M00027806:85         CH04MAL           700         2883         RTA22200249F.I.14.1.P.Seq         F         M00027717:75         CH04MAL           701         9784         RTA22200249F.c.12.1.P.Seq         F         M00027628:11         CH04MAL	Samuel	Samerana	kaan oo ka qoo ahaan oo ahaan		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
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698         6751         RTA22200248F.n.24.1.P.Seq         F         M00027604:710         CH04MAL           699         11619         RTA22200250F.d.17.1.P.Seq         F         M00027806:85         CH04MAL           700         2883         RTA22200249F.I.14.1.P.Seq         F         M00027717:75         CH04MAL           701         9784         RTA22200249F.c.12.1.P.Seq         F         M00027628:11         CH04MAL	English and the second	kanana yangan marangan marangan kananan kananan kananan kananan kananan kananan kananan kananan kananan kanana			arrangen arang panggan ara	and the second s
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Table 1

726         649655         RTA22200010F.o.22.1.P.Seq         F         M00056514:58         CH16COP           727         62016         RTA22200019F.k.05.1.P.Seq         F         M00054520:25         CH17COHLV           728         2783         RTA22200019F.k.18.1.P.Seq         F         M00054520:22         CH17COHLV           729         3876         RTA22200019F.c.03.1.P.Seq         F         M00043504:76         CH17COHLV           730         20036         RTA22200019F.f.11.1.P.Seq         F         M00054486:211         CH17COHLV           731         644032         RTA22200013F.e.06.1.P.Seq         F         M00043374:22         CH17COHLV           732         451636         RTA22200013F.e.01.1.P.Seq         F         M00024374:22         CH16COP           733         3428         RTA22200012F.p.22.1.P.Seq         F         M00056789:510         CH16COP           735         456506         RTA22200014F.p.23.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M00057035:39         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M00057039:16         CH16COP           740         4043		ble 1	050 14145	ODIENTATION	CLONETD	LIDDADY
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713         185432         RTA2220024F.m.02.1.P.Seq         F         M00005385:110         CH02COH           714         11456         RTA22200224F.m.02.1.P.Seq         F         M00005385:110         CH02COH           715         508892         RTA22200024F.f.14.1.P.Seq         F         M000056917:33         CH02COH           716         67         RTA22200224F.m.17.1.P.Seq         F         M00005695671         CH02COH           717         2636         RTA22200224F.m.16.1.P.Seq         F         M000058939:49         CH02COH           718         735028         RTA22200014F.e.08.1.P.Seq         F         M000058959:49         CH02COH           719         1924         RTA22200010F.m.24.1.P.Seq         F         M000056505:71         CH16COP           720         640116         RTA22200010F.m.24.1.P.Seq         F         M00005850:31         CH02COH           721         6546         RTA22200010F.p.03.1.P.Seq         F         M000068513:35         CH16COP           722         730866         RTA22200019F.b.03.1.P.Seq         F         M0005651:35         CH16COP           723         4829         RTA22200019F.b.03.1.P.Seq         F         M0005651:36         CH16COP           725         549934	CONTRACTOR OF THE CONTRACTOR O	European anno anno anno anno transcensione		λ <u></u>	egonouroagus gonourouro agente en en entergenoueuro en en en en en entere en	NOTES DE MINISTERNA DE CONTRA DE LA CONTRA DE
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721         6546         RTA22200225F.b.13.1.P.Seq         F         M00005420:31         CH02COH           722         730866         RTA22200010F.p.03.1.P.Seq         F         M000056515:35         CH16COP           723         4829         RTA22200224F.c.02.1.P.Seq         F         M0000485017:36         CH16COP           724         546632         RTA22200010F.o.12.1.P.Seq         F         M00054510:89         CH17COHLY           725         549934         RTA22200019F.i.19.1.P.Seq         F         M00054510:89         CH17COHLY           726         649655         RTA22200019F.k.18.1.P.Seq         F         M0005452:25         CH17COHLY           727         62016         RTA22200019F.k.18.1.P.Seq         F         M0005452:22         CH17COHLY           728         2783         RTA22200019F.e.03.1.P.Seq         F         M00043504:76         CH17COHLY           729         3876         RTA22200013F.e.06.1.P.Seq         F         M00043504:76         CH17COHLY           730         20036         RTA22200013F.e.01.1.P.Seq         F         M00043374:22         CH17COHLY           731         644032         RTA22200014F.e.01.1.P.Seq         F         M00022716:36         CH3COHY           732         451636	719	1924	RTA22200224F.n.16.1.P.Seq	F	M00005395:49	CH02COH
722         730866         RTA22200010F.p.03.1.P.Seq         F         M00056515:35         CH16COP           723         4829         RTA22200224F.c.02.1.P.Seq         F         M00004850:75         CH02COH           724         546632         RTA22200019F.o.12.1.P.Seq         F         M00056512:36         CH16COP           725         549934         RTA22200019F.i.19.1.P.Seq         F         M00056512:36         CH16COP           726         649655         RTA22200019F.b.05.1.P.Seq         F         M00054520:25         CH17COHLV           728         2783         RTA22200019F.b.05.1.P.Seq         F         M00043504:26         CH17COHLV           729         3876         RTA22200019F.b.05.1.P.Seq         F         M00043504:76         CH17COHLV           730         20036         RTA22200013F.e.06.1.P.Seq         F         M00043504:76         CH17COHLV           731         644032         RTA22200013F.e.06.1.P.Seq         F         M00056824:11         CH17COHLV           733         3428         RTA22200013F.e.06.1.P.Seq         F         M0005763:51         CH16COP           733         3428         RTA22200014F.p.22.1.P.Seq         F         M0005769:510         CH16COP           733         3428	720	640116	RTA22200010F.m.24.1.P.Seq		M00056503:711	CH16COP
723         4829         RTA22200224F.c.02.1.P.Seq         F         M00004850:75         CH02COH           724         546632         RTA22200010F.o.12.1.P.Seq         F         M00056512:36         CH16COP           725         549934         RTA22200010F.i.19.1.P.Seq         F         M00056514:38         CH16COP           726         649655         RTA22200019F.k.05.1.P.Seq         F         M00054520:25         CH17COHLV           727         62016         RTA22200019F.k.18.1.P.Seq         F         M00054520:25         CH17COHLV           728         2783         RTA22200019F.e.03.1.P.Seq         F         M00043504:76         CH17COHLV           730         2036         RTA22200019F.e.03.1.P.Seq         F         M00054524:22         CH17COHLV           731         644032         RTA22200013F.e.06.1.P.Seq         F         M00056824:51         CH16COP           732         451636         RTA22200013F.e.01.1.P.Seq         F         M000543374:22         CH17COHLV           733         3428         RTA22200014F.e.01.1.P.Seq         F         M000567369:510         CH16COP           735         456506         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           737         732712<	721	6546	RTA22200225F.b.13.1.P.Seq	F	M00005420:31	CH02COH
724         546632         RTA22200010F.o.12.1.P.Seq         F         M00056512:36         CH16COP           725         549934         RTA22200019F.i.19.1.P.Seq         F         M00056514:58         CH16COP           726         649655         RTA22200019F.c.022.1.P.Seq         F         M00056514:58         CH16COP           727         62016         RTA22200019F.k.05.1.P.Seq         F         M00054520:25         CH17COHLV           728         2783         RTA22200019F.e.03.1.P.Seq         F         M00054524:22         CH17COHLV           729         3876         RTA22200019F.e.03.1.P.Seq         F         M00054524:22         CH17COHLV           730         20036         RTA22200019F.e.03.1.P.Seq         F         M00056824:51         CH17COHLV           731         644032         RTA22200013F.e.06.1.P.Seq         F         M00056824:51         CH16COP           732         451636         RTA22200013F.e.01.1.P.Seq         F         M00043374:22         CH17COHLV           733         3428         RTA22200014F.e.02.P.Seq         F         M0004374:22         CH17COHLV           733         3456506         RTA22200014F.p.22.1.P.Seq         F         M0005709:10         CH16COP           735         456506 <td>722</td> <td>730866</td> <td>RTA22200010F.p.03.1.P.Seq</td> <td>F</td> <td>M00056515:35</td> <td>CH16COP</td>	722	730866	RTA22200010F.p.03.1.P.Seq	F	M00056515:35	CH16COP
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726         649655         RTA22200010F.o.22.1.P.Seq         F         M00056514:58         CH16COP           727         62016         RTA22200019F.k.05.1.P.Seq         F         M00054520:25         CH17COHLV           728         2783         RTA22200019F.k.18.1.P.Seq         F         M00054520:22         CH17COHLV           729         3876         RTA22200019F.f.11.1.P.Seq         F         M00043504:76         CH17COHLV           730         20036         RTA22200013F.e.06.1.P.Seq         F         M00056824:51         CH16COP           731         644032         RTA22200013F.e.06.1.P.Seq         F         M00043374:22         CH17COHLV           732         451636         RTA22200018F.o.24.1.P.Seq         F         M0002374:22         CH16COP           733         3428         RTA22200017F.p.22.1.P.Seq         F         M00056789:510         CH16COP           735         456506         RTA22200014F.p.23.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M00057035:39         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M0005709:9:16         CH16COP           740         4043	724	546632	RTA22200010F.o.12.1.P.Seq	F	M00056512:36	CH16COP
727         62016         RTA22200019F.k.05.1.P.Seq         F         M00054520:25         CH17COHL           728         2783         RTA22200019F.k.18.1.P.Seq         F         M00054524:22         CH17COHL           729         3876         RTA22200019F.e.03.1.P.Seq         F         M00054504:22         CH17COHL           730         20036         RTA22200019F.f.11.1.P.Seq         F         M00056824:51         CH16COP           731         644032         RTA22200013F.e.06.1.P.Seq         F         M00043374:22         CH17COHL           733         3428         RTA22200012F.p.22.1.P.Seq         F         M00022716:36         CH03MAH           734         643954         RTA22200012F.p.22.1.P.Seq         F         M00057150:310         CH16COP           735         456506         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M00057035:39         CH16COP           737         732712         RTA222000236F.c.13.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA222000236F.c.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043	725	549934	RTA22200019F.i.19.1.P.Seq	F	M00054510:89	CH17COHLV
728         2783         RTA22200019F.k.18.1.P.Seq         F         M00054524:22         CH17COHLV           729         3876         RTA22200019F.e.03.1.P.Seq         F         M00043504:76         CH17COHLV           730         20036         RTA22200013F.e.03.1.P.Seq         F         M00054846:211         CH16COP           731         644032         RTA22200013F.e.06.1.P.Seq         F         M0004374:22         CH17COHLV           732         451636         RTA22200013F.e.01.1.P.Seq         F         M00022716:36         CH03MAH           733         3428         RTA22200012F.p.22.1.P.Seq         F         M00022716:36         CH03MAH           734         643954         RTA22200014F.p.22.1.P.Seq         F         M0005703:310         CH16COP           735         456506         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M0002709:16         CH16COP           737         732712         RTA222000236F.c.13.1.P.Seq         F         M00025709:16         CH16COP           738         696         RTA22200235F.g.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043	726	649655	RTA22200010F.o.22.1.P.Seq	F	M00056514:58	CH16COP
729         3876         RTA22200019F.e.03.1.P.Seq         F         M00043504:76         CH17COHLV           730         20036         RTA22200019F.f.11.1.P.Seq         F         M00054486:211         CH17COHLV           731         644032         RTA22200013F.e.06.1.P.Seq         F         M00043374:22         CH17COHLV           732         451636         RTA22200018F.o.24.1.P.Seq         F         M00043374:22         CH17COHLV           733         3428         RTA22200012F.p.22.1.P.Seq         F         M00056789:510         CH16COP           734         643954         RTA22200012F.p.22.1.P.Seq         F         M0005789:510         CH16COP           735         456506         RTA22200014F.p.23.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           737         732712         RTA222000236F.c.13.1.P.Seq         F         M00057019:82         CH16COP           738         696         RTA22200024F.k.21.2.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200023F.g.13.1.P.Seq         F         M000586971:61         CH02COH           741         3639	727	62016	RTA22200019F.k.05.1.P.Seq	F	M00054520:25	CH17COHLV
729         3876         RTA22200019F.e.03.1.P.Seq         F         M00043504:76         CH17COHL           730         20036         RTA22200019F.f.11.P.Seq         F         M000564486:211         CH17COHL           731         644032         RTA22200018F.e.06.1.P.Seq         F         M00056824:51         CH16COP           732         451636         RTA22200018F.e.01.1.P.Seq         F         M0004374:22         CH17COHL           733         3428         RTA22200017F.e.01.1.P.Seq         F         M00056789:510         CH16COP           734         643954         RTA22200012F.p.22.1.P.Seq         F         M00056789:510         CH16COP           735         456506         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200024F.k.21.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200023F.g.13.1.P.Seq         F         M00057019:82         CH16COP           741         3639         <	728	2783	RTA22200019F.k.18.1.P.Seq	F	M00054524:22	CH17COHLV
731         644032         RTA22200013F.e.06.1.P.Seq         F         M00056824:51         CH16COP           732         451636         RTA22200018F.o.24.1.P.Seq         F         M00043374:22         CH17COHL           733         3428         RTA222000237F.e.01.1.P.Seq         F         M00056789:510         CH16COP           734         643954         RTA22200015F.p.22.1.P.Seq         F         M00057159:310         CH16COP           735         456506         RTA22200014F.p.23.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M00057035:39         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057099:16         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200024F.k.21.2.P.Seq         F         M00057019:82         CH16COP           741         3639         RTA22200228F.k.21.2.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200235F.g.13.1.P.Seq         F         M00022645:37         CH03MAH           743         1247	729	3876		F	M00043504:76	CH17COHLV
732         451636         RTA22200018F.o.24.1.P.Seq         F         M00043374:22         CH17COHLV           733         3428         RTA22200237F.e.01.1.P.Seq         F         M00022716:36         CH03MAH           734         643954         RTA22200012F.p.22.1.P.Seq         F         M00056789:510         CH16COP           735         456506         RTA22200015F.p.23.1.P.Seq         F         M00057035:39         CH16COP           736         449269         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M00057019:82         CH16COP           739         456528         RTA22200238F.k.21.2.P.Seq         F         M0006871:61         CH02COH           740         4043         RTA22200238F.k.21.2.P.Seq         F         M00022729:56         CH04MAL           741         3639         RTA22200236F.i.17.1.P.Seq         F         M000222472:511         CH03MAH           742         1024         RTA22200236F.i.13.2.P.Seq         F         M00022645:37         CH03MAH           744         4934	730	20036	RTA22200019F.f.11.1.P.Seq	F	M00054486:211	CH17COHLV
733         3428         RTA22200237F.e.01.1.P.Seq         F         M00022716:36         CH03MAH           734         643954         RTA22200012F.p.22.1.P.Seq         F         M00056789:510         CH16COP           735         456506         RTA22200015F.p.23.1.P.Seq         F         M00057150:310         CH16COP           736         449269         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200036F.c.13.1.P.Seq         F         M00057029:16         CH16COP           740         4043         RTA22200036F.c.13.1.P.Seq         F         M00057019:82         CH16COP           741         3639         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           743         1247         RTA22200236F.i.13.2.P.Seq         F         M00022645:37         CH03MAH           744         4934         R	731	644032	RTA22200013F.e.06.1.P.Seq	F	M00056824:51	CH16COP
734         643954         RTA22200012F.p.22.1.P.Seq         F         M00056789:510         CH16COP           735         456506         RTA22200015F.p.23.1.P.Seq         F         M00057150:310         CH16COP           736         449269         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M0002579:512         CH03MAH           739         456528         RTA22200014F.n.13.1.P.Seq         F         M000057019:82         CH16COP           740         4043         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH04MAL           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200237F.f.07.1.P.Seq         F         M0005699.311         CH16COP           745         901 <td< td=""><td>732</td><td>451636</td><td>RTA22200018F.o.24.1.P.Seq</td><td>F</td><td>M00043374:22</td><td>CH17COHLV</td></td<>	732	451636	RTA22200018F.o.24.1.P.Seq	F	M00043374:22	CH17COHLV
734         643954         RTA22200012F.p.22.1.P.Seq         F         M00056789:510         CH16COP           735         456506         RTA22200015F.p.23.1.P.Seq         F         M00057150:310         CH16COP           736         449269         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M0002599:512         CH03MAH           739         456528         RTA22200014F.n.13.1.P.Seq         F         M000057019:82         CH16COP           740         4043         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200236F.i.13.2.P.Seq         F         M0005699.311         CH16COP           745         901 <td< td=""><td>733</td><td>3428</td><td>RTA22200237F.e.01.1.P.Seq</td><td>F</td><td>M00022716:36</td><td>CH03MAH</td></td<>	733	3428	RTA22200237F.e.01.1.P.Seq	F	M00022716:36	CH03MAH
735         456506         RTA22200015F.p.23.1.P.Seq         F         M00057150:310         CH16COP           736         449269         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200218F.k.21.2.P.Seq         F         M00057019:82         CH16COP           741         3639         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200236F.i.17.1.P.Seq         F         M00022472:51         CH03MAH           743         1247         RTA22200236F.i.13.2.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200238F.i.13.2.P.Seq         F         M0002645:37         CH03MAH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00056990:311         CH16COP           747         725825         RTA		643954	RTA22200012F.p.22.1.P.Seg	F	M00056789:510	CH16COP
736         449269         RTA22200014F.p.17.1.P.Seq         F         M00057035:39         CH16COP           737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M00022598:512         CH03MAH           739         456528         RTA22200014F.n.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200228F.k.21.2.P.Seq         F         M00057019:82         CH16COP           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200236F.i.17.1.P.Seq         F         M00022472:511         CH03MAH           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M0006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808	735	456506		F	M00057150:310	CH16COP
737         732712         RTA22200014F.p.03.1.P.Seq         F         M00057029:16         CH16COP           738         696         RTA22200236F.c.13.1.P.Seq         F         M00022598:512         CH03MAH           739         456528         RTA22200014F.n.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200244F.p.09.1.P.Seq         F         M00022645:37         CH03MAH           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00002645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M00006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.04.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056925:37         CH16COP           750         551907	CONTRACTOR CONTRACTOR CONTRACTOR					LANGERTON OF THE PROPERTY AND ADDRESS OF THE PARTY OF THE
738         696         RTA22200236F.c.13.1.P.Seq         F         M00022598:512         CH03MAH           739         456528         RTA22200014F.n.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200244F.p.09.1.P.Seq         F         M00022645:37         CH03MAH           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M0006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M0005694:44         CH03MAH           746         452726         RTA22200014F.i.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RT		Anna and a second secon	and the contraction of the contr	F	M00057029:16	CH16COP
739         456528         RTA22200014F.n.13.1.P.Seq         F         M00057019:82         CH16COP           740         4043         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200244F.p.09.1.P.Seq         F         M00027229:56         CH04MAL           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M0006846:43         CH02COH           745         901         RTA222000237F.f.07.1.P.Seq         F         M0005694:34         CH03MAH           746         452726         RTA22200014F.I.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.I.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH16COP           752         7098         R		3		F	M00022598:512	CH03MAH
740         4043         RTA22200228F.k.21.2.P.Seq         F         M00006871:61         CH02COH           741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200244F.p.09.1.P.Seq         F         M00027229:56         CH04MAL           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M00006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00056924:44         CH03MAH           746         452726         RTA22200014F.l.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA222000014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           751         551527         RTA222000237F.n.22.1.P.Seq         F         M00056998:58         CH16COP           752         7098 <t< td=""><td></td><td>\$</td><td></td><td>F</td><td>M00057019:82</td><td>CH16COP</td></t<>		\$		F	M00057019:82	CH16COP
741         3639         RTA22200235F.g.13.1.P.Seq         F         M00022472:511         CH03MAH           742         1024         RTA22200244F.p.09.1.P.Seq         F         M00027229:56         CH04MAL           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M00006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00022724:44         CH03MAH           746         452726         RTA22200014F.I.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00056998:58         CH16COP           754         554812 <t< td=""><td></td><td>4043</td><td></td><td>F</td><td>CONTRACTOR OF THE CONTRACTOR O</td><td></td></t<>		4043		F	CONTRACTOR OF THE CONTRACTOR O	
742         1024         RTA22200244F.p.09.1.P.Seq         F         M00027229:56         CH04MAL           743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M00006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00022724:44         CH03MAH           746         452726         RTA22200014F.i.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200023F.n.22.1.P.Seq         F         M00056998:58         CH16COP           753         4589         RTA22200225F.k.24.1.P.Seq         F         M0005699:38         CH16COP           755         3114         RT		}		F		
743         1247         RTA22200236F.i.17.1.P.Seq         F         M00022645:37         CH03MAH           744         4934         RTA22200228F.i.13.2.P.Seq         F         M00006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00022724:44         CH03MAH           746         452726         RTA22200014F.i.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00056998:58         CH16COP           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00056969:38         CH16COP           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         <		\$~~~~~~~~~~~~~		F		
744         4934         RTA22200228F.i.13.2.P.Seq         F         M00006846:43         CH02COH           745         901         RTA22200237F.f.07.1.P.Seq         F         M00022724:44         CH03MAH           746         452726         RTA22200014F.i.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH15CON           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		Barrer				
745         901         RTA22200237F.f.07.1.P.Seq         F         M00022724:44         CH03MAH           746         452726         RTA22200014F.i.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH15CON           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M00056969:38         CH16COP           754         554812         RTA22200023F.g.01.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH	Contraction of the Contraction o	(magazaran managazaran managazaran managazaran managazaran managazaran managazaran managazaran managazaran man	aran arang ara			
746         452726         RTA22200014F.I.04.1.P.Seq         F         M00057003:29         CH16COP           747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH15CON           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M00056969:38         CH16COP           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		}		X	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
747         725825         RTA22200014F.i.23.1.P.Seq         F         M00056990:311         CH16COP           748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH15CON           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M00056969:38         CH16COP           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		Samuel Sa		,		
748         456808         RTA22200014F.a.19.1.P.Seq         F         M00056924:26         CH16COP           749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH15CON           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M0005501:55         CH02COH           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		å	······································	<u> </u>		
749         729295         RTA22200014F.a.21.1.P.Seq         F         M00056925:37         CH16COP           750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH15CON           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M0005501:55         CH02COH           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH						
750         551907         RTA22200005F.a.18.1.P.Seq         F         M00055850:63         CH15CON           751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M00005501:55         CH02COH           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		iyaanaa aa a				
751         551527         RTA22200014F.k.03.1.P.Seq         F         M00056998:58         CH16COP           752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M00005501:55         CH02COH           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		5.,, <u>,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,</u>				
752         7098         RTA22200237F.n.22.1.P.Seq         F         M00022831:87         CH03MAH           753         4589         RTA22200225F.k.24.1.P.Seq         F         M00005501:55         CH02COH           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		Samuel and the second second				
753         4589         RTA22200225F.k.24.1.P.Seq         F         M00005501:55         CH02COH           754         554812         RTA22200014F.g.21.1.P.Seq         F         M00056969:38         CH16COP           755         3114         RTA22200232F.g.01.1.P.Seq         F         M00022015:611         CH03MAH		i de la composição de l	an ann an airm an an an airm an airm an an an an an airm an airm an an airm an an an an an an an an an airm ai	Common a servicio de common de servicio de common de como de c	agranus de agranus conservir de Adrigan, en vers mes del Frida escribir de Adrifes com escabilità del	AND THE PROPERTY OF THE PROPERTY OF THE PARTY OF THE PART
754 554812 RTA22200014F.g.21.1.P.Seq F M00056969:38 CH16COP 755 3114 RTA22200232F.g.01.1.P.Seq F M00022015:611 CH03MAH		·			***************************************	
755 3114 RTA22200232F.g.01.1.P.Seq F M00022015:611 CH03MAH			uuun meeruun aan araan aan aan aan aan aan aan aan			
		<u></u>				
- 756 : 6031 : RTAZZZUGZZZEM 13.1 P.SAG : - : MOODO6/31:43 : CHO2COH	756	6031	RTA22200232F.g.01.1.F.Seq	F	M000022013.011	CH02COH

Table 1

Та	ble 1				
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759	3428	RTA22200237F.d.24.1.P.Seq	F	M00022716:36	CH03MAH
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762	234605	RTA22200009F.c.17.2.P.Seq	F	M00042515:110	CH16COP
763	2224	RTA22200232F.g.14.1.P.Seq	F	M00022049:27	CH03MAH
764	185642	RTA22200243F.a.07.1.P.Seq	F	M00027076:67	CH04MAL
765	649655	RTA22200007F.d.16.1.P.Seq	F	M00056171:811	CH15CON
Samuel and the same of the sam	2854	RTA22200231F.o.23.1.P.Seq	F	M00007998:67	CH03MAH
766	453470	RTA22200007F.j.08.1.P.Seq	F	M00056224:58	CH15CON
767	<u> </u>	RTA22200027F.k.14.1.P.Seq	F	M00006704:76	CH02COH
768	11012	RTA22200005F.c.23.1.P.Seq	F	M00055869:16	CH15CON
769	535208	RTA22200003F.C.23.1.F.3eq	F	M00056994:33	CH16COP
770	448606	RTA22200014F.J.14.1.F.Seq	F	M00027172:33	CH04MAL
771	12304		F	M00006989:54	CH02COH
772	2756	RTA22200229F.h.17.1.P.Seq	F	M00055871:65	CH15CON
773	367	RTA22200005F.d.04.1.P.Seq	F	M000033871.03	CH02COH
774	11351	RTA22200229F.c.09.1.P.Seq	F F	M00006888:75	CH02COH
775	6858	RTA22200228F.n.03.2.P.Seq	F	M00006889:711	CH04MAL
776	7750	RTA22200241F.g.01.1.P.Seq	3		CH03MAH
777	6923	RTA22200232F.h.12.1.P.Seq	<u> </u>	M00022060:69	
778	11552	RTA22200232F.p.03.1.P.Seq	F	M00022163:38	CH03MAH
779	12448	RTA22200243F.c.10.1.P.Seq	Į F	M00027090:58	CH04MAL
780	2944	RTA22200227F.m.11.1.P.Seq	F	M00006731:22	CH02COH
781	10342	RTA22200231F.I.03.1.P.Seq	F	M00007981:24	CH03MAH
782	9026	RTA22200222F.o.20.1.P.Seq	F	M00004304:47	CH01COH
783	10342	RTA22200238F.m.23.1.P.Seq	F	M00022997:38	CH03MAH
784	6455	RTA22200235F.f.08.1.P.Seq	F	M00022465:65	CH03MAH
785	6455	RTA22200232F.b.15.1.P.Seq	F	M00021886:66	CH03MAH
786	3416	RTA22200227F.k.16.1.P.Seq	F	M00006705:79	CH02COH
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788	2889	RTA22200228F.n.19.2.P.Seq	F	M00006894:13	CH02COH
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790	14390	RTA22200244F.i.10.1.P.Seq	F	M00027193:35	CH04MAL
791	661	RTA22200241F.a.15.1.P.Seq	F	M00026856:311	CH04MAL
792	452992	RTA22200013F.e.07.1.P.Seq	F	M00056826:212	CH16COP
793	1943	RTA22200235F.k.21.2.P.Seq		M00022528:812	CH03MAH
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795	5482	RTA22200225F.p.15.1.P.Seq		M00005565:65	CH02COH
796	650493	RTA22200007F.c.10.1.P.Seq		M00056156:112	CH15CON
797	640318	RTA22200007F.b.04.1.P.Seq		M00056145:24	CH15CON
798	646309	RTA22200011F.p.23.1.P.Seq	F	M00056661:55	CH16COP
799	4316	RTA22200248F.e.15.1.P.Seq	F	M00027543:77	CH04MAL
800	449701	RTA22200012F.d.12.1.P.Seq		M00056695:89	CH16COP
801	560367	RTA22200013F.e.03.1.P.Seq		M00056824:310	CH16COP
802	9997	RTA22200238F.o.18.1.P.Seq	en antigente alle extreme in commence de la commenc	M00023015:42	CH03MAH
803	649106	RTA22200012F.f.03.1.P.Seq		M00056707:52	CH16COP
804	461835	RTA22200007F.b.18.1.P.Seq		M00056150:312	CH15CON
805	640590	RTA22200003F.m.09.1.P.Sec		M00055703:35	CH15CON
806	648340	RTA22200007F.a.08.1.P.Seq	and the second s	M00056139:54	CH15CON
807	554812	RTA22200014F.g.21.2.P.Seq	and designation of the second	M00056969:38	CH16COP
808	447035	RTA22200007F.m.06.1.P.Sec		M00056252:511	CH15CON
809	1208	RTA22200238F.g.03.1.P.Seq		M00022908:83	CH03MAH
810	3114	RTA22200232F.f.24.1.P.Seq	and make the second	M00022015:611	CH03MAH
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Table 1

Та	ble 1				
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813	450323	RTA22200015F.o.17.1.P.Seq	F	M00057141:42	CH16COP
814	11567	RTA22200232F.f.16.1.P.Seq	F	M00022010:81	CH03MAH
815	11567	RTA22200232F.c.14.1.P.Seq	F	M00021915:510	CH03MAH
816	6660	RTA22200222F.j.01.1.P.Seq	F	M00003986:712	CH01COH
817	9026	RTA22200222F.n.12.1.P.Seq	F	M00004198:111	CH01COH
	185539	RTA22200243F.b.09.1.P.Seq	F	M00027084:810	CH04MAL
818	<u> </u>	RTA22200223F.d.01.1.P.Seq	F	M00001537:610	CH01COH
819	3224	RTA22200240F.I.12.1.P.Seq	F	M00026804:43	CH04MAL
820	95700	RTA22200240F.I.12.1.F.Seq	F	M00004167:411	CH01COH
821	4439	RTA22200222F.II.01.1.F.Seq	F	M00022667:511	CH03MAH
822	3428		F	M00022681:56	CH03MAH
823	1456	RTA22200236F.p.05.1.P.Seq	F	M000022001:50	CH02COH
824	11343	RTA22200228F.g.03.1.P.Seq	F	M00056746:46	CH16COP
825	729206	RTA22200012F.k.20.1.P.Seq	F F	M00057147:11	CH16COP
826	558371	RTA22200015F.p.13.1.P.Seq		M00057147.11	CH16COP
827	451589	RTA22200011F.m.21.1.P.Seq	<u> </u>		CH16COP
828	404475	RTA22200012F.f.10.1.P.Seq	<u> </u>	M00056708:36	and the second s
829	734582	RTA22200017F.e.07.1.P.Seq	<u> </u>	M00057334:61	CH16COP
830	729779	RTA22200011F.i.11.1.P.Seq	F	M00056596:52	CH16COP
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832	449269	RTA22200014F.p.17.2.P.Seq	F	M00057035:39	CH16COP
833	4609	RTA22200227F.a.11.1.P.Seq	F	M00006583:83	CH02COH
834	640318	RTA22200011F.i.13.1.P.Seq	İ F	M00056596:56	CH16COP
835	729851	RTA22200011F.j.22.1.P.Seq	F	M00056610:812	CH16COP
836	11028	RTA22200230F.a.15.1.P.Seq	F	M00007098:35	CH02COH
837	643924	RTA22200008F.e.17.1.P.Seq	F	M00056345:53	CH15CON
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839	11286	RTA22200230F.k.07.1.P.Seq	F	M00007194:56	CH02COH
840	185651	RTA22200248F.g.16.1.P.Seq	F	M00027556:710	CH04MAL
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842	728408	RTA22200014F.p.13.2.P.Seq	F	M00057033:69	CH16COP
843	646309	RTA22200007F.e.15.1.P.Seq	F	M00056184:83	CH15CON
844	405073	RTA22200003F.m.14.1.P.Seq	F	M00055704:47	CH15CON
845	185489	RTA22200241F.j.24.1.P.Seq	F	M00026926:35	CH04MAL
846	447326	RTA22200005F.i.15.1.P.Seq	F	M00055909:58	CH15CON
847	11006	RTA22200227F.c.15.1.P.Seq	i F	M00006610:48	CH02COH
848	6863	RTA22200227F.a.23.1.P.Seq		M00006588:86	CH02COH
849	11351	RTA22200225F.p.12.1.P.Seq	······································	M00005563:212	CH02COH
850	401553	RTA22200005F.f.13.1.P.Seq		M00055884:21	CH15CON
851	504513	RTA22200007F.h.19.1.P.Seq		M00056213:14	CH15CON
852	645979	RTA22200012F.g.11.1.P.Seq	and province and our conservations are a series and a series and a series are a series and a series are a series and a series are a ser	M00056713:78	CH16COP
\$ AND	6923	RTA222000121 .g.11.11 .coq		M00008023:412	
853	1924	RTA22200239F.I.18.1.P.Seq		M00007034:26	CH02COH
854	washing and the second	RTA22200230F.f.02.1.P.Seq	<u></u>	M00007136:35	CH02COH
855	5838	RTA22200230F.f.24.1.P.Seq	and the second s	M00007146:611	<u>เรื่องแบบแบบแบบเทคเหมองทางเพลเหม และเคมองการการการกา</u>
856	2062	RTA22200003F.b.06.1.P.Seq	<u></u>	M00055581:21	CH15CON
857	447388	RTA22200003F.5.06.1.P.Seq		M00026865:711	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
858	12419			M00023663711	akan marana ana ana ana ana ana ana ana ana an
859	3224	RTA22200222F.c.24.1.P.Seq	market and the second s	M00007174:46	CH02COH
860	5474	RTA22200230F.i.23.1.P.Seq		M00007174:40	CH02COH
861	3522	RTA22200227F.f.23.1.P.Seq		M00006030.43	CH16COP
862	731785	RTA22200012F.o.21.1.P.Seq		M00036777.07	CH02COH
863	3765	RTA22200228F.d.10.1.P.Seq	The second secon	M00056664:76	CH16COP
864	640323	RTA22200012F.a.10.1.P.Sec	F F	1010000004.70	1 0,110001

Table 1

	ble 1				LIDDADY
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866	448029	RTA22200012F.n.03.1.P.Seq	F	M00056764:58	CH16COP
867	650476	RTA22200005F.h.07.1.P.Seq	F	M00055893:35	CH15CON
868	640525	RTA22200007F.f.12.1.P.Seq	F	M00056192:82	CH15CON
869	390124	RTA22200007F.n.09.1.P.Seq	F	M00056267:52	CH15CON
870	464029	RTA22200023F.p.21.1.P.Seq	F	M00055137:34	CH17COHLV
871	468109	RTA22200005F.e.18.1.P.Seq	F	M00055880:612	CH15CON
872	21669	RTA22200023F.h.20.1.P.Seq	F	M00055064:512	CH17COHLV
873	651088	RTA22200007F.j.07.2.P.Seq	F	M00056224:210	CH15CON
874	2737	RTA22200023F.d.15.1.P.Seq	F	M00055039:71	CH17COHLV
875	556421	RTA22200005F.e.14.1.P.Seq	F	M00055879:44	CH15CON
876	452245	RTA22200023F.c.23.1.P.Seq	F	M00055034:81	CH17COHLV
877	447539	RTA22200005F.e.22.1.P.Seq	F	M00055882:19	CH15CON
878	546642	RTA22200014F.o.20.2.P.Seq	F	M00057029:312	CH16COP
879	236368	RTA22200006F.j.23.2.P.Seq	F	M00056077:56	CH15CON
880	644523	RTA22200012F.b.12.1.P.Seq	F	M00056673:56	CH16COP
881	729173	RTA22200012F.k.02.1.P.Seq	F	M00056739:411	CH16COP
882	8315	RTA22200231F.a.06.1.P.Seq	F	M00007927:31	CH03MAH
883	450463	RTA22200007F.i.11.2.P.Seq	F	M00056218:73	CH15CON
884	650856	RTA22200012F.o.13.1.P.Seq	F	M00056775:31	CH16COP
885	648109	RTA22200005F.i.14.1.P.Seq	F	M00055909:71	CH15CON
886	726644	RTA22200013F.d.19.1.P.Seq	F	M00056822:711	CH16COP
887	727224	RTA22200013F.h.19.1.P.Seq	F	M00056859:412	CH16COP
888	557906	RTA22200023F.h.18.1.P.Seq	F	M00055063:71	CH17COHLV
889	502683	RTA22200007F.e.22.1.P.Seq	F	M00056185:46	CH15CON
890	728408	RTA22200011F.i.06.1.P.Seq	F	M00056594:310	CH16COP
891	647952	RTA22200005F.j.11.1.P.Seq	F	M00055919:16	CH15CON
892	639991	RTA22200009F.I.23.2.P.Seq	F	M00042846:49	CH16COP
893	735346	RTA22200016F.m.21.1.P.Seq	F	M00057266:44	CH16COP
894	102655	RTA22200007F.i.23.2.P.Seq	F	M00056222:62	CH15CON
895	553629	RTA22200012F.o.03.1.P.Seq	F	M00056772:58	CH16COP
896	1609	RTA22200226F.i.23.1.P.Seq	F	M00005708:27	CH02COH
897	641884	RTA22200005F.I.16.1.P.Seq	F	M00055937:32	CH15CON
898	648872	RTA22200007F.p.10.1.P.Seq	adriani and a second	M00056286:112	CH15CON
899	644242	RTA22200005F.i.20.1.P.Seq	F	M00055911:56	CH15CON
900	63559	RTA22200244F.f.14.1.P.Seq	F	M00027183:21	CH04MAL
901	550108	RTA22200012F.o.24.1.P.Seq	F	M00056779:810	CH16COP
902	374306	RTA22200011F.n.05.1.P.Seq		M00056639:52	CH16COP
903	5838	RTA22200226F.c.19.1.P.Seq		M00005621:88	CH02COH
904	645530	RTA22200006F.k.07.2.P.Seq		M00056079:67	CH15CON
905	649732	RTA22200012F.b.05.1.P.Seq	<u></u>	M00056669:77	CH16COP
906	649143	RTA22200007F.p.01.1.P.Seq		M00056283:52	CH15CON
907	7571	RTA22200225F.h.24.1.P.Seq		M00005481:36	CH02COH
5		RTA22200236F.j.09.1.P.Seq	F	M00022651:31	CH03MAH
908	4572 2147	RTA22200230F.j.09.1.F.Seq		M00007006:11	CH02COH
909	management and control of the second	RTA22200008F.h.10.1.P.Seq		M00056475:62	CH15CON
910	462659 727723	RTA222000081.11.10.111.364		M00056761:65	CH16COP
911		RTA22200012F.III.21.1.F.Seq		M00005657:112	and province and resemble and recommendation of the second
912	2636	RTA22200226F.g.21.1.P.Seq		M00055931:51	CH15CON
913	500959	RTA22200005F.K.21.1.F.Seq		M00007954:24	CH03MAH
914	3428	RTA22200231F.1.07.1.P.Seq		M00056592:47	CH16COP
915	734929			M00030392.47	
916	453592	RTA22200001F.g.19.1.P.Seq RTA22200026F.d.21.1.P.Seq		M00055423:810	na kanana na manana n
917	15414		and the same and t	M00055542:19	Land and the commence of the c
918	648959	RTA22200002F.m.22.1.P.Sec	<u> 11                                  </u>	1000000042.19	

Table 1

Та	ble 1				LIBBABY
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922	649719	RTA22200006F.m.21.2.P.Seq	F	M00056098:81	CH15CON
923	562805	RTA22200023F.d.08.1.P.Seq	F	M00055037:510	CH17COHLV
924	452204	RTA22200001F.p.24.1.P.Seq	F	M00054918:311	CH15CON
925	549178	RTA22200023F.d.22.1.P.Seq	F	M00055041:52	CH17COHLV
926	639177	RTA22200015F.n.24.1.P.Seq	F	M00057134:710	CH16COP
927	562550	RTA22200022F.i.05.1.P.Seq	F	M00054941:74	CH17COHLV
928	561807	RTA22200004F.I.07.1.P.Seq	F	M00055808:711	CH15CON
929	641373	RTA22200006F.m.23.2.P.Seq	F	M00056099:79	CH15CON
930	514418	RTA22200022F.a.15.1.P.Seq	F	M00054870:49	CH17COHLV
931	567078	RTA22200006F.b.18.2.P.Seq	F	M00056007:111	CH15CON
932	643061	RTA22200006F.n.01.2.P.Seq	F	M00056099:811	CH15CON
933	549160	RTA22200020F.i.09.1.P.Seq	F	M00054638:38	CH17COHLV
934	449269	RTA22200009F.e.07.1.P.Seq	F	M00042770:212	CH16COP
935	453082	RTA22200001F.e.03.1.P.Seq	F	M00042564:811	CH15CON
936	418135	RTA22200026F.a.09.1.P.Seq	F	M00055406:45	CH17COHLV
937	2783	RTA22200023F.b.23.1.P.Seq	F	M00055027:47	CH17COHLV
938	549435	RTA22200007F.j.06.2.P.Seq	F	M00056223:73	CH15CON
939	446614	RTA22200009F.i.13.2.P.Seq	F	M00042816:64	CH16COP
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940	454380	RTA22200012F.p.04.11F.Seq	F	M00056815:21	CH16COP
941	450914	RTA22200013F.c.05.1.P.Seq	F	M00056812:58	CH16COP
943	736860	RTA22200016F.p.14.1.P.Seq	F	M00057283:56	CH16COP
943	727224	RTA222000101.p.14.11004	F	M00056592:64	CH16COP
945	644242	RTA22200006F.a.19.2.P.Seq	F	M00056000:612	
945	562550	RTA222000001.a.10.2.1.00q	F	M00055002:54	CH17COHLV
947	649148	RTA22200007F.j.24.2.P.Seq	F	M00056231:79	CH15CON
948	375889	RTA22200023F.c.13.1.P.Seq	F	M00055032:16	CH17COHLV
949	449437	RTA22200009F.e.03.1.P.Seq	F	M00042767:42	CH16COP
950	449044	RTA22200006F.b.07.2.P.Seq	F	M00056003:56	CH15CON
951	555318	RTA22200022F.e.11.1.P.Seq	F	M00054903:312	
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953	11567	RTA22200231F.g.22.1.P.Seq		M00007965:710	£
954	3522	RTA22200229F.j.18.1.P.Seq	F F	M00007007:85	CH02COH
955	456528	RTA22200016F.p.05.1.P.Seq	F	M00057280:16	CH16COP
	639142	RTA22200013F.m.02.1.P.Seq		M00056887:68	CH16COP
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957	554742	RTA222000011.e.10.1.10cq		M00055405:85	CH17COHLV
958	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	RTA222000261 .a.03.1.1 .0cq		M00042848:712	
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961	551527			M00056925:37	CH16COP
962	729295	RTA22200014F.a.21.2.P.Seq RTA22200022F.m.17.1.P.Seq		M00054975:34	CH17COHLV
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964	648996	RTA22200004F.d.22.1.P.Seq		M00033708.23	CH15CON
965	447126	RTA22200001F.e.18.1.P.Seq		M00042373:17 M00056916:22	CH16COP
966	730866	RTA22200013F.p.20.1.P.Seq	<u></u>	M00054692:41	CH17COHLV
967	420686	RTA22200020F.o.02.1.P.Seq		M00057304:51	CH16COP
968	451753	RTA22200017F.b.07.1.P.Seq		M00054743:52	CH17COHLV
969	451380	RTA22200021F.d.12.2.P.Seq	F F	M00055689:67	CH15CON
970	645530	RTA22200003F.I.11.1.P.Seq		M00057266:712	
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972	562835	RTA22200015F.n.20.1.P.Seq	F	M00057134:11	

Table 1

	ble 1				LIDDADY
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975	728779	RTA22200017F.c.19.1.P.Seq	F	M00057318:29	CH16COP
976	414739	RTA22200002F.n.19.1.P.Seq	F	M00055547:410	CH15CON
977	551514	RTA22200022F.k.05.1.P.Seq	F	M00054953:710	CH17COHLV
978	550107	RTA22200023F.p.02.1.P.Seq	F	M00055130:71	CH17COHLV
979	726786	RTA22200017F.a.10.1.P.Seq	F	M00057291:26	CH16COP
980	456747	RTA22200001F.f.23.1.P.Seq	F	M00042700:11	CH15CON
981	562550	RTA22200023F.n.13.1.P.Seq	F	M00055111:43	CH17COHLV
European Commence	549722	RTA22200025F.p.13.1.P.Seq	F	M00055560:66	CH15CON
982	640525	RTA22200004F.i.10.1.P.Seq	F	M00055796:510	CH15CON
983		RTA22200009F.f.23.1.P.Seq	F	M00042787:59	CH16COP
984	455542		F	M00057270:54	CH16COP
985	9436	RTA22200016F.n.12.1.P.Seq	F	M00057270:54 M00055755:83	CH15CON
986	380284	RTA22200004F.c.17.1.P.Seq	F	M00053755.85	CH17COHLV
987	556260	RTA22200020F.p.08.1.P.Seq		5	CH16COP
988	650476	RTA22200013F.p.21.1.P.Seq	<u> </u>	M00056916:64	Ž.
989	554500	RTA22200006F.n.14.2.P.Seq	F	M00056103:812	CH15CON
990	422375	RTA22200016F.n.16.1.P.Seq	F	M00057272:23	CH16COP
991	456528	RTA22200019F.I.23.1.P.Seq	F	M00054535:89	CH17COHLV
992	644190	RTA22200009F.m.23.1.P.Seq	F	M00042853:73	CH16COP
993	554080	RTA22200014F.o.19.3.P.Seq	F	M00057028:49	CH16COP
994	546705	RTA22200021F.i.06.3.P.Seq	F	M00054781:24	CH17COHLV
995	558337	RTA22200021F.a.02.2.P.Seq	F	M00054720:611	
996	449269	RTA22200014F.p.17.3.P.Seq	F	M00057035:39	CH16COP
997	645799	RTA22200014F.d.22.2.P.Seq	F	M00056952:84	CH16COP
998	456506	RTA22200001F.I.10.1.P.Seq	F	M00042889:19	CH15CON
999	218416	RTA22200026F.c.09.1.P.Seq	F	M00055414:111	CH17COHLV
1000	455820	RTA22200023F.m.12.1.P.Seq	F	M00055097:76	CH17COHLV
1001	554703	RTA22200024F.p.19.1.P.Seq	F	M00055262:211	CH17COHLV
1002	650204	RTA22200008F.c.16.1.P.Seq	F	M00056313:67	CH15CON
1003	456808	RTA22200014F.a.19.2.P.Seq	F	M00056924:26	CH16COP
1004	420686	RTA22200004F.n.01.1.P.Seq	F	M00055821:16	CH15CON
1005	378373	RTA22200023F.n.20.1.P.Seq	F	M00055114:111	CH17COHLV
1006	463824	RTA22200004F.a.22.1.P.Seq	F	M00055740:69	CH15CON
1007	24939	RTA22200016F.i.15.1.P.Seq	F	M00057229:66	CH16COP
1007	556561	RTA22200022F.n.08.1.P.Seq	F	M00054981:36	CH17COHLV
<u> </u>	380406	RTA222000221 III.00.11 .00q	F	M00042890:38	CH15CON
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1013	5830		<u> </u>	M00056909:49	CH16COP
1014	539955	RTA22200013F.o.22.1.P.Seq	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M00055553:37	CH15CON
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1021	559806	RTA22200022F.o.02.1.P.Seq		M00054987:32	CH17COHLV
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1026	641683	RTA22200003F.k.24.1.P.Seq	F	M00055685:51	CH15CON

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1027	648934	RTA22200003F.a.23.1.P.Seq	F	M00055578:89	CH15CON
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1029	377094	RTA22200021F.d.05.2.P.Seq	F	M00054741:510	CH17COHLV
1030	449617	RTA22200026F.c.15.1.P.Seq	F	M00055415:811	CH17COHLV
1031	978	RTA22200004F.m.17.1.P.Seq	F	M00055820:58	CH15CON
1032	607430	RTA22200024F.n.10.1.P.Seq	F	M00055247:111	CH17COHLV
1033	641837	RTA22200003F.b.20.1.P.Seq	F	M00055586:65	CH15CON
1034	449750	RTA22200021F.h.08.3.P.Seq	F	M00054773:112	CH17COHLV
1035	646780	RTA22200004F.m.23.1.P.Seq	F	M00055820:710	CH15CON
1036	546642	RTA22200014F.o.20.3.P.Seq	F	M00057029:312	CH16COP
1037	642906	RTA22200014F.i.13.2.P.Seq	F	M00056985:35	CH16COP
1037	552879	RTA22200024F.g.10.1.P.Seq	F	M00055196:19	CH17COHLV
- Same and the sam	<u></u>	RTA2220003F.d.03.1.P.Seq	F	M00055602:710	CH15CON
1039	644205 506744	RTA222000031 .d.03.111 .064	F	M00054841:27	CH17COHLV
1040	San and the san an	RTA22200024F.c.03.1.P.Seq	F	M00055154:64	CH17COHLV
1041	557797	RTA22200024F.b.18.1.P.Seq	F	M00055792:57	CH15CON
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1043	462659	RTA22200001F.i.16.1.P.Seq	F	M00042732.79	CH15CON
1044	645633	RTA22200008F.a.01.1.P.Seq	F F	M00055495:63	CH15CON
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1046	454343	RTA22200016F.d.13.1.P.Seq		M00057181.40	CH17COHLV
1047	386543	RTA22200024F.g.20.1.P.Seq	<u> </u>	describerations de la company de la comp	CH15CON
1048	446404	RTA22200003F.b.10.1.P.Seq	<u> </u>	M00055582:54	CH16COP
1049	456528	RTA22200014F.n.13.2.P.Seq	<u> </u>	M00057019:82	CH16COP
1050	456528	RTA22200015F.I.08.1.P.Seq	<u> </u>	M00057118:32	
1051	452781	RTA22200004F.o.03.1.P.Seq	Į F	M00055827:36	CH15CON
1052	551671	RTA22200021F.b.20.2.P.Seq	F	M00054729:51	CH17COHLV
1053	644242	RTA22200004F.h.08.1.P.Seq	<u> </u>	M00055791:45	CH15CON
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1055	450429	RTA22200004F.e.09.1.P.Seq	<u> </u>	M00055771:41	CH15CON
1056	533588	RTA22200001F.k.13.1.P.Seq	ļ F	M00042881:38	CH15CON
1057	553877	RTA22200004F.f.03.1.P.Seq	<u> </u>	M00055775:210	
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1062	736816	RTA22200015F.b.20.1.P.Seq	F	M00057052:79	CH16COP
1063	403632	RTA22200020F.b.08.1.P.Seq	and a superior contract of the	M00054575:31	CH17COHLV
1064	390124	RTA22200021F.n.17.2.P.Seq	F	M00054849:811	and processes and the second s
1065	390124	RTA22200016F.i.16.1.P.Seq	F	M00057230:412	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
1066	422687	RTA22200015F.e.14.1.P.Seq		M00057072:52	CH16COP
1067	394413	RTA22200001F.k.11.1.P.Seq		M00042750:29	CH15CON
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1069	453079	RTA22200004F.e.11.1.P.Seq		M00055771:111	
1070	463824	RTA22200001F.j.03.1.P.Seq	F	M00042735:17	CH15CON
1071	736595	RTA22200016F.d.16.1.P.Seq	F.	M00057182:211	,,,5,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
1072	102655	RTA22200004F.m.02.1.P.Sec		M00055816:61	CH15CON
1073	448606	RTA22200014F.j.14.2.P.Seq		M00056994:33	CH16COP
1074	504513	RTA22200004F.n.12.1.P.Seq		M00055823:411	
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1076	530883	RTA22200016F.d.08.1.P.Seq	F	M00057180:811	
1077	447126	RTA22200004F.o.18.1.P.Seq		M00055832:512	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
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1080	549320	RTA22200020F.g.16.1.P.Seq		M00054621:411	CH17COHLV
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	CLUSTER	SEQ NAME	ORIENTATION	M00055306:79	CH17COHL
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1082	450791	RTA22200018F.k.03.1.P.Seq	F	M00043338:13	CH17COHL
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1091	97507	RTA22200025F.i.22.1.P.Seq	F	M00055364:51	CH17COH
1092	556216	RTA22200025F.k.07.1.P.Seq	F	M00055373:410	CH17COH
1093	185401	RTA22200250F.d.22.1.P.Seq	F	M00027808:710	CH04MA
1094	3758	RTA22200248F.o.08.1.P.Seq	F	M00027605:55	CH04MA
1095	95700	RTA22200248F.I.15.1.P.Seq	F	M00027588:16	CH04MA
1096	2478	RTA22200250F.d.12.1.P.Seq	F	M00027803:810	CH04MA
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1098	185652	RTA22200250F.c.19.1.P.Seq	F	M00027786:21	CH04MA
1098	55798	RTA22200250F.a.04.1.P.Seq	F	M00027757:26	CH04MA
	3	RTA222002301.a.04.111.3cq	F	M00056529:89	CH16CO
1100	5078	RTA22200011F.a.19.1.F.3eq	F	M00027681:42	CH04MA
1101	9784	Lancard Company of the Company of th	F	M00027001:42 M00022216:46	CH03MA
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1103	11606	RTA22200234F.e.19.1.P.Seq	F F	M00022221:46	CH03MA
1104	2245	RTA22200234F.e.21.1.P.Seq	F	M00022221.46 M00056537:19	CH16CO
1105	551172	RTA22200011F.b.19.1.P.Seq			ž
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1107	6317	RTA22200224F.e.13.1.P.Seq	F	M00004971:74	CH02CO
1108	2478	RTA22200234F.f.21.1.P.Seq	F	M00022235:311	CH03MA
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1118	554785	RTA22200019F.h.01.1.P.Seq	F	M00054499:511	CH17COF
1119	551235	RTA22200019F.e.14.1.P.Seq	F	M00043506:89	CH17COH
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****		RTA22200005F.b.15.1.P.Seq	F	M00055856:67	CH15CC
1124	550267		F	M00022697:412	<del>valerium anno anno anno anno anno anno anno ann</del>
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1126	451794	RTA22200005F.c.13.1.P.Seq			and for the second seco
1127	5744	RTA22200227F.I.15.1.P.Seq	F	M00006719:512	<u> </u>
1128	3516	RTA22200228F.n.13.2.P.Seq	F	M00006892:69	CH02CC
1129	730555	RTA22200014F.j.23.1.P.Seq	<u>F</u>	M00056997:89	CH16CC
1130	3085	RTA22200237F.n.05.1.P.Seq	<u> </u>	M00022829:86	CH03MA
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RTA22200005F.a.16.1.P.Seq

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M00027571:311

M00055849:48

CH04MAL

CH15CON

Table 1

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1135	646248	RTA22200005F.a.14.1.P.Seq	F	M00055848:86	CH15CON
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1138	6923	RTA22200237F.h.10.1.P.Seq	F	M00022741:211	CH03MAH
1139	6923	RTA22200237F.h.16.1.P.Seq	F	M00022745:37	CH03MAH
1140	901	RTA22200237F.o.03.1.P.Seq	F	M00022831:34	CH03MAH
1141	901	RTA22200237F.n.23.1.P.Seq	F	M00022831:19	CH03MAH
1142	367	RTA22200236F.h.18.1.P.Seq	F	M00022641:310	CH03MAH
1143	4043	RTA22200228F.I.13.2.P.Seq	F	M00006873:21	CH02COH
1144	3299	RTA22200236F.h.19.1.P.Seq	F	M00022641:56	CH03MAH
1145	11881	RTA22200238F.e.15.1.P.Seq	F	M00022899:39	CH03MAH
1146	9113	RTA22200230F.I.04.1.P.Seq	F	M00007204:712	CH02COH
1147	185460	RTA22200243F.p.24.1.P.Seq	F	M00027165:611	CH04MAL
1148	185716	RTA22200241F.d.03.1.P.Seq	F	M00026873:511	CH04MAL
1149	5753	RTA22200227F.o.22.1.P.Seq	F	M00006756:68	CH02COH
1150	24939	RTA22200012F.e.06.1.P.Seq	F	M00056701:58	CH16COP
1151	649684	RTA22200007F.a.14.1.P.Seq	F	M00056140:87	CH15CON
1151	642109	RTA22200017.d.11	F	M00056717:34	CH16COP
1152	15035	RTA2220007F.I.11.1.P.Seq	F	M00056246:23	CH15CON
\$	649354	RTA22200007F.a.15.1.P.Seq	F	M00056140:57	CH15CON
1154	4465	RTA222000071.a.13.111.06q	F	M00006811:412	CH02COH
1155		RTA222002287.e.19.1.F.Seq	F.	M00056144:39	CH15CON
1156	647952	RTA22200007F.b.02.1.F.Seq	F	M00055990:25	CH15CON
1157	455601	RTA22200005F.p.18.1.F.Seq	F	M00055984:32	CH15CON
1158	641901	RTA22200003F.b.07.1.F.3eq	F	M00042466:86	CH16COP
1159	446878	RTA22200009F.b.21.2.F.Seq	F	M00072168:11	CH03MAH
1160	7436 2245	RTA22200232F.ii.08.1.F.Seq	F	M00007166:56	CH02COH
1161	3531	RTA22200230F.i.03.FF06q	F	M00006746:26	CH02COH
1162	9625	RTA222002271.0.01.1.1.1.Seq	F F	M00023520:77	CH04MAL
1163	727489	RTA22200012F.I.19.1.P.Seq	F	M00056754:14	CH16COP
1164 1165	159925	RTA22200240F.j.14.1.P.Seq	F	M00023428:43	CH04MAL
\$	645210	RTA22200012F.g.17.1.P.Seq	<u></u>	M00056715:54	CH16COP
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1167	8375	RTA222002331.d.24.1.10eq	F	M00007982:611	CH03MAH
1168		RTA22200231F.i.07:1.F.Seq		M00007129:68	CH02COH
1169	4319	RTA22200230F.e.02.1.F.Seq	<u></u>	M00007992:78	CH03MAH
1170	4045	RTA222002311.11.07.1.11.0eq		M00026848:711	CH04MAL
1171	185642	RTA22200238F.I.11.1.P.Seq	F	M00022974:410	CH03MAH
1172	7436	RTA222002361.111.1.00q	and for the second contract of the second con	M00006746:26	CH02COH
1173	3531	RTA222002271:11:24:111:00q	<u></u>	M00056715:58	CH16COP
1174	644776	RTA222000121 .g.21.111 .Geq		M00022892:77	CH03MAH
1175	8354	RTA222002361.d.13.1.1beq	F	M00027111:84	CH04MAL
1176	2099	RTA22200243F.I.03.1.F.Seq	F	M00056624:85	CH16COP
1177	449956	3		M00055723:55	CH15CON
1178	649106	RTA22200003F.o.09.1.P.Seq RTA22200005F.i.09.1.P.Seq	† F	M00055908:512	CH15CON
1179	452414	3		M000556616:110	Samuel and the second s
1180	732712	RTA22200011F.k.21.1.P.Seq		M00030010:116	CH04MAL
1181	185562	RTA22200241F.m.01.1.P.Sec RTA22200225F.i.17.1.P.Seq		M00020335:04 M00005485:19	CH02COH
1182	3516	1		M00003403.13	CH04MAL
1183	185562	RTA22200241F.I.24.1.P.Seq		M00020938.04 M00026854:57	CH04MAL
1184	185460	RTA22200241F.a.08.1.P.Seq		M00020034:37	CH03MAH
1185	10947	RTA22200237F.I.16.1.P.Seq		M00056169:66	CH15CON
1186	452856	RTA22200007F.d.13.1.P.Seq		M00057127:77	CH16COP
1187	558767	RTA22200015F.m.17.1.P.Sec	Ladjuurus varanna maaran maa maa maa maa maa maa maa maa maa m	M00057127.77	CH15CON
1188	15035	RTA22200008F.e.22.1.P.Sec	L <u>í</u>	101000000400.10	1 011100014

Table 1

	ble 1				
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1190	7082	RTA22200235F.o.24.2.P.Seq	F	M00022565:15	CH03MAH
1191	452523	RTA22200013F.j.07.1.P.Seq	F	M00056868:59	CH16COP
1192	3242	RTA22200230F.a.01.1.P.Seq	F	M00007092:63	CH02COH
1193	6660	RTA22200222F.i.24.1.P.Seq	F	M00003986:712	CH01COH
1194	547	RTA22200244F.j.02.1.P.Seq	F	M00027197:77	CH04MAL
1195	121213	RTA22200011F.i.23.1.P.Seq	F	M00056599:411	CH16COP
1196	4378	RTA22200226F.p.15.1.P.Seq	F	M00005830:410	CH02COH
1197	185554	RTA22200244F.m.17.1.P.Seq	F	M00027217:73	CH04MAL
1198	185482	RTA22200241F.d.04.1.P.Seq	F	M00026873:28	CH04MAL
1199	185715	RTA22200240F.I.14.1.P.Seq	F	M00026805:24	CH04MAL
1200	66017	RTA22200243F.d.10.1.P.Seq	F	M00027097:711	CH04MAL
1201	403111	RTA22200007F.j.14.1.P.Seq	F	M00056226:612	CH15CON
1202	3224	RTA22200222F.o.18.1.P.Seq	F	M00004296:711	CH01COH
1203	966	RTA22200238F.k.11.1.P.Seq	F	M00022961:211	CH03MAH
1204	3639	RTA22200235F.j.05.2.P.Seq	F	M00022509:26	CH03MAH
1205	5388	RTA22200243F.k.17.1.P.Seq	F	M00027139:36	CH04MAL
1206	3299	RTA22200238F.b.05.1.P.Seq	F	M00022872:25	СН03МАН
1207	23760	RTA22200241F.n.13.1.P.Seq	F	M00026949:810	CH04MAL
1208	729384	RTA22200012F.n.06.1.P.Seq	F	M00056765:512	CH16COP
1209	46559	RTA22200016F.g.16.1.P.Seq	F	M00057215:22	CH16COP
1210	449750	RTA22200022F.n.05.1.P.Seg	F	M00054980:32	CH17COHLV
1211	735936	RTA22200011F.i.02.1.P.Seq	F	M00056593:55	CH16COP
1212	607430	RTA22200005F.o.04.1.P.Seq	F	M00055971:57	CH15CON
1213	452856	RTA22200007F.g.23.1.P.Seq	F	M00056205:41	CH15CON
1214	557903	RTA22200016F.g.07.1.P.Seq	F	M00057211:67	CH16COP
1215	453112	RTA22200001F.g.01.1.P.Seq	F	M00042700:43	CH15CON
1216	645900	RTA22200006F.i.24.2.P.Seq	F	M00056067:48	CH15CON
1217	415114	RTA22200002F.m.20.1.P.Seg	F	M00055542:31	CH15CON
1218	418763	RTA22200004F.I.11.1.P.Seq	F	M00055811:18	CH15CON
1219	2245	RTA22200230F.a.12.1.P.Seq	F	M00007097:24	CH02COH
1220	403668	RTA22200012F.o.18.1.P.Seq	F	M00056777:33	CH16COP
1221	15427	RTA22200020F.m.08.1.P.Seq	F	M00054677:42	CH17COHLV
1222	555714	RTA22200020F.n.23.1.P.Seq	F :	M00054691:55	CH17COHLV
1223	555830	RTA22200022F.d.19.1.P.Seq	F	M00054899:67	CH17COHLV
1224	4620	RTA22200231F.e.13.1.P.Seq	F	M00007951:15	CH03MAH
1225	171511	RTA22200012F.p.21.1.P.Seq	F	M00056789:34	CH16COP
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1227	447501	RTA22200002F.e.01.1.P.Seq	F	M00055453:51	CH15CON
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1237	645305	RTA22200001F.p.17.1.P.Seq	F	M00054917:69	CH15CON
1238	463487	RTA222000017.p.17.11.P.Seq	F	M00056184:48	CH15CON
1239	11131	RTA222000077.e.12.1.F.Seq	F	M000007105:312	CH02COH
1240	561807	RTA22200200F.k.18.1.P.Seq	F	M00007103.312	CH17COHLV
1241	452800	RTA22200020F.n.03.1.P.Seq	F	M00055104:212	CH17COHLV
1241	372960	RTA22200023F.n.03.1.F.Seq	F	M00056782:26	CH16COP
1474	012300	1 174222000 121 .P.07 . 1.F . 364		19100000102.20	OHIOOOP

Ta	ble 1			C C C C C C C C C C C C C C C C C C C	LIDDADY
SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
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1245	9113	RTA22200226F.i.12.1.P.Seq	F	M00005704:410	CH02COH
1246	630259	RTA22200004F.k.14.1.P.Seq	F	M00055805:410	CH15CON
1247	3516	RTA22200226F.o.20.1.P.Seq	F	M00005819:211	CH02COH
1248	447494	RTA22200004F.j.14.1.P.Seq	F	M00055802:72	CH15CON
1249	554500	RTA22200019F.n.10.1.P.Seq	F	M00054547:59	CH17COHLV
1250	639662	RTA22200001F.p.19.1.P.Seq	F	M00054917:412	CH15CON
1251	421	RTA22200009F.o.23.1.P.Seq	F	M00042869:56	CH16COP
1252	736014	RTA22200016F.o.19.1.P.Seq	F	M00057277:39	CH16COP
1253	643061	RTA22200006F.m.24.2.P.Seq	F	M00056099:811	CH15CON
1254	9113	RTA22200229F.m.02.1.P.Seq	F	M00007035:56	CH02COH
1255	650856	RTA22200007F.i.19.2.P.Seq	F	M00056221:55	CH15CON
1256	476223	RTA22200009F.g.09.1.P.Seq	F	M00042792:64	CH16COP
1257	737088	RTA22200011F.f.01.1.P.Seq	F	M00056564:59	CH16COP
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1267	552614	RTA22200008F.e.13.1.P.Seq	F	M00056344:73	CH15CON
1268	452523	RTA22200007F.g.15.1.P.Seq	F	M00056203:810	CH15CON
1269	446789	RTA22200004F.I.01.1.P.Seq	F	M00055807:710	CH15CON
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Same and the same	403111	RTA22200005F.m.11.1.P.Seq		M00055946:77	CH15CON
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1283	403111	RTA222000071.iii.22.111.3000		M00056066:87	CH15CON
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1290	455492	RTA22200001F.j.08.1.P.Seq		M00057079:59	
1291	639667	RTA22200015F.f.11.1.P.Seq	and an investment and a second	M00057079:55 M00055181:51	CH17COHL\
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1294	561485	RTA22200026F.d.14.1.P.Sec	and the second s	M00055528:66	
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1296	451401	RTA22200018F.h.16.1.P.Sec	η F	17.04	CITICONE

Table 1

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1305	401553	RTA22200004F.e.16.1.P.Seq	F	M00055771:67	CH15CON
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1312	730555	RTA22200014F.j.23.2.P.Seq	F	M00056997:89	CH16COP
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1319	644721	RTA22200004F.p.13.1.P.Seq	F	M00055839:69	CH15CON
1320	453202	RTA22200025F.n.08.2.P.Seq	F	M00055390:48	CH17COHLV
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1322	641988	RTA22200002F.c.20.1.P.Seq	F	M00055446:26	CH15CON
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1349	648959	RTA22200002F.g.21.1.F.Seq	F	***************************************	<del> </del>
1350			anno como a como a como a como como como co	M00042537:85	CH15CON
1330	380291	RTA22200001F.b.12.1.P.Seq	F	M00042537:85	CH15CON

Table 1

	DIE 1				
dramatic colors become production	CLUSTER	SEQ NAME	ORIENTATION	CONTRACTOR OF THE PROPERTY OF A SECURIOR STATE OF THE PROPERTY	LIBRARY
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1355	452833	RTA22200018F.d.24.1.P.Seq	F.	M00042520:69	CH17COHLV
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1359	549829	RTA22200020F.f.14.1.P.Seq	F	M00054609:86	CH17COHLV
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1364	727181	RTA22200011F.d.23.1.P.Seq	F	M00056555:69	CH16COP
1365	454050	RTA22200011F.c.06.1.P.Seq	F	M00056541:18	CH16COP
1366	725994	RTA22200011F.b.07.1.P.Seq	F	M00056534:411	CH16COP
1367	1495	RTA22200234F.j.11.1.P.Seq	F	M00022273:19	CH03MAH
1368	5665	RTA22200233F.n.01.1.P.Seq	F	M00021654:14	CH03MAH
1369	5665	RTA22200233F.m.24.1.P.Seq	F	M00021654:14	CH03MAH
1370	646146	RTA22200010F.i.03.1.P.Seq	F	M00056421:612	CH16COP
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1378	555928	RTA22200010F.o.23.1.P.Seq	F	M00056514:71	CH16COP
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1392	380477	RTA22200007F.d.03.1.P.Seq	F	M00056162:46	CH15CON
1393	3299	RTA22200238F.a.22.1.P.Seq	F	M00022861:210	CH03MAH
1394	448853	RTA22200009F.b.12.2.P.Seq	F	M00042463:69	CH16COP
1395	736701	RTA22200012F.h.21.1.P.Seq	F	M00056723:410	CH16COP
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Table 1

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1409	5156	RTA22200225F.m.12.1.P.Seq	F	M00005516:86	CH02COH
1410	728408	RTA22200012F.h.05.1.P.Seq	F	M00056718:72	CH16COP
1411	73812	RTA22200242F.m.20.1.P.Seq	F	M00027054:23	CH04MAL
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1414	5240	RTA22200230F.i.16.1.P.Seq	F	M00007172:33	CH02COH
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1445	734582	RTA22200013F.n.18.1.P.Seq	<b>F</b>	M00056816:110	CH16COP
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1447	463487 558710	RTA22200001F.e.05.1.P.Seq	F	M00042565:83	CH15CON
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Table 1

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1472	450255	RTA22200021F.j.13.3.P.Seq	F	M00054810:82	CH17COHLV
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1482	428005	RTA22200021F.c.03.2.P.Seq	<u> </u>	M00054731:312	CH17COHLV
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1504	450255	RTA22200018F.c.16.1.P.Seq	F	M00042455:411	CH17COHLV
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1509	726307	RTA22200010F.o.16.1.P.Seq	F	M00056512:312	S
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1511	3524	RTA22200227F.f.19.1.P.Seq	F	M00006650:13	CH02COH
1512	8112	RTA22200237F.m.13.1.P.Seq	0.\$10.00.00.00.00.00.00.00.00.00.00.00.00.0	M00022823:43	CH03MAH
1012	1 0112	1 177222002071 .III. 10. 1.1° .OEG	<u> </u>	17100022020.70	

Table 1

	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
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1515	2676	RTA22200227F.k.01.1.P.Seq	F	M00006695:88	CH02COH
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1519	8336	RTA22200235F.f.24.1.P.Seq	<u> </u>	M00022470:71	CH03MAH
1520	185542	RTA22200248F.h.05.1.P.Seq	<u> </u>	M00027564:43	CH04MAL
1521	448046	RTA22200014F.d.13.1.P.Seq	<u> </u>	M00056951:69	CH16COP
1522	185422	RTA22200242F.j.15.1.P.Seq	<u> </u>	M00027034:411	CH04MAL
1523	650448	RTA22200005F.c.17.1.P.Seq	<u>F</u>	M00055868:66	CH15CON
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1533	2622	RTA22200248F.i.16.1.P.Seq	F	M00027573:69	CH04MAL
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1535	19205	RTA22200244F.b.14.1.P.Seq	F	M00027171:27	CH04MAL
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1538	779	RTA22200238F.f.01.1.P.Seq	F	M00022901:511	CH03MAH
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1540	5289	RTA22200235F.m.20.2.P.Seq	F	M00022550:24	CH03MAH
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1549	401849	RTA22200003F.p.02.1.P.Seq	F	M00055726:412	CH15CON
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1554	3656	RTA22200227F.f.01.1.P.Seq	F	M00006641:83	CH02COH
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1562	452531	RTA22200009F.J.08.2.P.Seq	F	M00042842:58	CH16COP
1563	559057	RTA222000313.00.23 .Seq	F	M00042042:36	CH17COHLV
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	ble 1				
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1568	172013	RTA22200002F.c.05.1.P.Seq	F	M00055441:42	CH15CON
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1570	446531	RTA22200001F.c.11.1.P.Seq	F	M00042547:411	CH15CON
1571	639352	RTA22200008F.a.09.1.P.Seq	F	M00056293:79	CH15CON
1572	642604	RTA22200004F.e.12.1.P.Seq	F	M00055771:65	CH15CON
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£		RTA22200013F.II.06.1.P.Seq	F	Company of the Compan	december of the contract of th
1581	734554		F	M00057024:88	CH16COP
1582	418008	RTA22200020F.e.16.1.P.Seq		M00054601:810	CH17COHLV
1583	558614	RTA22200024F.k.16.1.P.Seq	<u> </u>	M00055227:89	CH17COHLV
1584	452245	RTA22200025F.I.16.1.P.Seq	E	M00055381:59	CH17COHLV
1585	449891	RTA22200019F.b.13.1.P.Seq	<u> </u>	M00043397:22	CH17COHLV
1586	547916	RTA22200011F.a.18.1.P.Seq	F	M00056529:612	CH16COP
1587	6162	RTA22200233F.I.19.1.P.Seq	F	M00021628:211	CH03MAH
1588	6162	RTA22200234F.f.14.1.P.Seq	F	M00022234:45	CH03MAH
1589	4809	RTA22200224F.d.21.1.P.Seq	F	M00004868:412	CH02COH
1590	3926	RTA22200242F.d.18.1.P.Seq	F	M00026993:86	CH04MAL
1591	185693	RTA22200248F.d.20.1.P.Seq	F	M00027532:32	CH04MAL
1592	641683	RTA22200007F.m.16.1.P.Seq	F	M00056256:811	CH15CON
1593	11351	RTA22200226F.o.18.1.P.Seq	F	M00005818:29	CH02COH
1594	650864	RTA22200008F.g.16.1.P.Seq	F	M00056467:57	CH15CON
1595	460445	RTA22200001F.i.11.1.P.Seq	F	M00042728:57	CH15CON
1596	447669	RTA22200011F.e.16.1.P.Seq	F	M00056561:48	CH16COP
1597	227936	RTA22200016F.m.13.1.P.Seq	F	M00057259:810	CH16COP
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1599	650195	RTA22200004F.a.04.1.P.Seq	F	M00055734:85	CH15CON
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1603	4045	RTA22200016F.1.21.1.P.Seq	F F	M00057208:36	Samonno como anterior a como como a como como a como como com
1604	4045 447669	RTA22200232F.g.23.1.P.Seq	F	M00056628:61	CH16COP
1605	11351	RTA22200011F.I.23.1.P.Seq	F		\$ 000000000000000000000000000000000000
	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			M00007121:78	CH02COH
1607	648931	RTA22200002F.I.16.1.P.Seq	F	M00055530:27	CH15CON
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1613	729851	RTA22200013F.h.15.1.P.Seq	F	M00056858:23	CH16COP
1614	2512	RTA22200235F.c.11.1.P.Seq	F	M00022430:410	CH03MAH
1615	452704	RTA22200008F.f.21.1.P.Seq	F	M00056456:61	CH15CON
1616	4589	RTA22200230F.c.15.1.P.Seq	F	M00007117:83	CH02COH
1617	4727	RTA22200233F.e.16.1.P.Seq	F	M00008059:28	CH03MAH
1618	454380	RTA22200014F.d.23.1.P.Seq	F	M00056953:16	CH16COP
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Table 1

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1623	448511	RTA22200023F.b.05.1.P.Seq	F	M00055021:411	CH17COHLV
1624	335	RTA22200012F.I.18.1.P.Seq	F	M00056753:110	CH16COP
1625	561382	RTA22200026F.c.05.1.P.Seq	F	M00055413:27	CH17COHLV
1626	3447	RTA22200227F.k.15.1.P.Seq	F	M00006705:19	CH02COH
1627	639896	RTA22200012F.c.15.1.P.Seq	F	M00056684:611	CH16COP
1628	1353	RTA22200241F.e.20.1.P.Seq	F	M00026890:42	CH04MAL
1629	3031	RTA22200229F.i.23.1.P.Seq	F	M00006996:610	CH02COH
1630	557928	RTA22200021F.g.03.3.P.Seq	F	M00054765:35	CH17COHLV
1631	4727	RTA22200237F.n.09.1.P.Seq	F	M00022829:810	CH03MAH
1632	4046	RTA22200226F.g.16.1.P.Seq	F	M00005655:68	CH02COH
1633	10882	RTA22200241F.b.11.1.P.Seq	F	M00026861:19	CH04MAL
1634	646283	RTA22200007F.c.22.1.P.Seq	F	M00056161:74	CH15CON
1635	646283	RTA22200002F.d.05.1.P.Seg	F	M00055448:57	CH15CON
1636	139516	RTA22200242F.e.16.1.P.Seq	F	M00027000:65	CH04MAL
1637	6184	RTA22200230F.e.13.1.P.Seq	F	M00007131:11	CH02COH
1638	6184	RTA22200228F.c.12.1.P.Seq	F	M00006795:13	CH02COH
1639	454653	RTA22200013F.e.17.1.P.Seq	F	M00056830:72	CH16COP
1640	3309	RTA22200232F.c.11.1.P.Seq	F	M00021911:83	CH03MAH
1641	1037	RTA22200248F.e.02.1.P.Seq	F	M00027535:58	CH04MAL
1642	450665	RTA22200002F.g.06.1.P.Seq	F	M00055491:78	CH15CON
1643	726307	RTA22200016F.b.16.1.P.Seq	F	M00057167:512	CH16COP
1644	447669	RTA22200001F.b.14.1.P.Seq	F	M00042538:56	CH15CON
1645	639651	RTA22200003F.b.15.1.P.Seq	F	M00055584:76	CH15CON
1646	736860	RTA22200015F.b.11.1.P.Seq	F	M00057049:76	CH16COP
1647	553705	RTA22200025F.j.22.1.P.Seq	F	M00055371:61	CH17COHLV
1648	451375	RTA22200018F.j.20.1.P.Seq	F	M00043336:58	CH17COHLV
1649	204862	RTA22200006F.h.24.2.P.Seq	F	M00056058:84	CH15CON
1650	530883	RTA22200011F.o.17.1.P.Seq	F	M00056652:64	CH16COP
1651	447539	RTA22200001F.e.22.1.P.Seq	F	M00042575:41	CH15CON
1652	455096	RTA22200020F.m.23.1.P.Seq	F	M00054680:211	}
1653	8336	RTA22200235F.g.01.1.P.Seq	F	M00022470:71	CH03MAH
1654	449142	RTA22200006F.f.09.2.P.Seq	F	M00056041:34	CH15CON
1655	557401	RTA22200003F.j.04.1.P.Seq	F	M00055668:27	CH15CON
1656	418763	RTA22200002F.e.07.1.P.Seq	F	M00055454:75	CH15CON
1657	17649	RTA22200238F.b.23.1.P.Seq	F	M00022876:48	CH03MAH
1658	2078	RTA22200238F.h.11.1.P.Seq	F	M00022928:63	СНОЗМАН
1659	640370	RTA22200010F.g.14.1.P.Seq	F	M00056410:54	CH16COP
1660	449269	RTA22200005F.h.22.1.P.Seq	F	M00055906:410	CH15CON
1661	639029	RTA22200006F.o.10.2.P.Seq	F	M00056114:36	CH15CON
1662	448677	RTA22200023F.I.07.1.P.Seq	F	M00055088:59	CH17COHLV
1663	349	RTA22200231.i.07111.i.ocq	F	M000000000:03	CH03MAH
1664	447494	RTA22200002F.e.05.1.P.Seq	F	M00055454:811	CH15CON
1665	551433	RTA222000021 .e.03.1.1 .Seq	F	M00053434:871	CH17COHLV
1666	414739	RTA222000217 .k. 12.3.F. Seq	F	M00034810.411	CH17COHLV
1667	640525	RTA22200019F.1.07.1.F.Seq	F	M00045508.31	CH15CON
1668	640525	RTA22200002F.m.24.1.P.Seq	F	M00055543:35	CH15CON
1669	233108	RTA22200002F.m.24.1.F.Seq	F	M00055560:212	CH15CON CH15CON
1670	643594	RTA22200002F.B.10.1.P.Seq		M00055687:61	CH15CON
1670	1642	RTA22200003F.1.06.1.P.Seq	F	M00007931:110	
1672	643804	RTA2220023TF.a.18.1.P.Seq	F	M00007931:110	CH03MAH CH15CON
1673	449701	RTA22200003F.h.09.1.P.Seq	F	M00056723:311	CH16COP
1673	185695	RTA22200012F.II.22.1.F.Seq	F	M00036723.311 M00026986:810	CH04MAL
10/4	100030	1 1 7 2 2 2 0 0 2 4 2 1 . U.Z 1 . I . F . O C Q	ı	141000020300.0 TO	CHOTIMAL

Table 1

	ble 1				
SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	THE PROPERTY AND ADDRESS OF THE PROPERTY OF TH	LIBRARY
1675	555830	RTA22200003F.k.13.1.P.Seq	F	M00055679:17	CH15CON
1676	227936	RTA22200010F.n.11.1.P.Seq	F	M00056505:47	CH16COP
1677	1609	RTA22200226F.g.09.1.P.Seq	F	M00005650:16	CH02COH
1678	643938	RTA22200005F.o.22.1.P.Seq	F	M00055980:110	CH15CON
1679	3656	RTA22200227F.e.24.1.P.Seq	F	M00006641:83	CH02COH
1680	16576	RTA22200248F.e.07.1.P.Seq	F	M00027536:712	CH04MAL
1681	9784	RTA22200243F.c.16.1.P.Seq	F	M00027093:28	CH04MAL
1682	2557	RTA22200228F.p.12.2.P.Seq	F	M00006917:48	CH02COH
1683	4620	RTA22200237F.o.15.1.P.Seq	F	M00022834:71	CH03MAH
1684	43642	RTA22200021F.p.23.2.P.Seq	F	M00054865:84	CH17COHLV
1685	555103	RTA22200016F.I.08.1.P.Seq	F	M00057249:36	CH16COP
1686	643341	RTA22200002F.g.16.1.P.Seq	F	M00055495:72	CH15CON
1687	185531	RTA22200248F.I.12.1.P.Seq	F	M00027588:31	CH04MAL
1688	4045	RTA22200224F.b.17.1.P.Seq	F	M000027000:01	CH02COH
1689	400258	RTA22200211F.e.10.1.P.Seq	F	M00056557:42	CH16COP
1690	96618	RTA22200011F.e.10.1.F.Seq	F	M00030337.42 M00027581:51	CH04MAL
Secretaria de la constantida del constantida de la constantida de la constantida de la constantida del constantida de la constantida del constantida d	646060	RTA22200246F.j.22.1.F.Seq	F	M00027381.31	CH15CON
1691	burramanan and an and an	***************************************	F	the complete for the control of the	TO THE PARTY OF TH
1692	5665	RTA22200232F.a.17.1.P.Seq	F	M00021854:57	CH03MAH
1693	149265	RTA22200241F.o.03.1.P.Seq		M00026951:43	CH04MAL
1694	727314	RTA22200012F.g.14.1.P.Seq	<u> </u>	M00056714:86	CH16COP
1695	736349	RTA22200014F.d.03.1.P.Seq	<u> </u>	M00056947:69	CH16COP
1696	648931	RTA22200006F.k.13.2.P.Seq	<u>F</u>	M00056081:25	CH15CON
1697	553881	RTA22200003F.m.10.1.P.Seq	F	M00055703:79	CH15CON
1698	7444	RTA22200235F.d.02.1.P.Seq	F	M00022440:41	CH03MAH
1699	150	RTA22200235F.p.14.2.P.Seq	F	M00022571:411	CH03MAH
1700	2889	RTA22200228F.p.09.2.P.Seq	F	M00006917:15	CH02COH
1701	730670	RTA22200013F.a.09.1.P.Seq	F	M00056793:87	CH16COP
1702	560984	RTA22200021F.n.20.2.P.Seq	F	M00054851:53	CH17COHLV
1703	453708	RTA22200026F.e.10.1.P.Seq	F	M00055425:35	CH17COHLV
1704	48977	RTA22200024F.p.08.1.P.Seq	F	M00055259:64	CH17COHLV
1705	547916	RTA22200008F.g.01.1.P.Seq	F	M00056459:37	CH15CON
1706	547916	RTA22200008F.f.24.1.P.Seg	F	M00056459:37	CH15CON
1707	97507	RTA22200023F.p.11.1.P.Seq	F	M00055134:82	CH17COHLV
1708	735966	RTA22200012F.n.08.1.P.Seq	F	M00056766:110	CH16COP
1709	35	RTA22200012F.m.20.1.P.Seq	F	M00056760:14	CH16COP
1710	650195	RTA22200002F.m.11.1.P.Seq	F	M00055536:63	CH15CON
1711	639705	RTA22200002F.i.11.1.P.Seq	F	M00055510:27	CH15CON
1712	185465	RTA22200013F.c.17.1.P.Seq	F	M00056817:33	CH16COP
1713	378525	RTA22200015F.h.03.1.P.Seq	F	M00057091:33	CH16COP
1714	2889	RTA22200228F.p.10.2.P.Seq	F	M00006917:14	CH02COH
1715	557686	RTA22200025F.f.19.1.P.Seq	F	M00055334:89	CH17COHLV
1716	735786	RTA22200010F.h.02.1.P.Seq	F	M00056414:15	CH16COP
1717	455145	RTA22200003F.f.10.1.P.Seq	F	M00055636:610	CH15CON
1718	639667	RTA22200002F.k.13.1.P.Seq	F	M00055525:27	CH15CON
1719	446913	RTA22200001F.m.21.1.P.Seq	F	M00042905:310	CH15CON
1720	402494	RTA22200014F.c.09.2.P.Seq	F	M00056941:52	CH16COP
1721	734256	RTA22200014F.I.11.1.P.Seq	F	M00057005:43	CH16COP
1722	734256	RTA22200012F.g.05.1.P.Seq	F	M00057005.45	CH16COP
1723	559362	RTA22200012F.g.05.1.F.Seq	F	M00055115:55	CH17COHLV
- Bremanne arrenn manuer anni	harmon, no reconstruction and representation of the second	RTA22200023F.II.21.1.P.Seq	F	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Anomy was now as the green was now a series of the series
1724	639651			M00055597:58	CH15CON
1725	419774	RTA22200008F.g.11.1.P.Seq	<u> </u>	M00056466:52	CH15CON
1726	555318	RTA22200025F.a.17.1.P.Seq	F	M00055271:35	CH17COHLV
1727	449956	RTA22200010F.b.08.1.P.Seq	<u> </u>	M00056359:111	CH16COP
1728	558427	RTA22200015F.p.12.1.P.Seq	F	M00057146:39	CH16COP

Table 1

	ble 1			<del></del>	
SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	MANAGEMENT AND	LIBRARY
1729	7531	RTA22200225F.j.13.1.P.Seq	<u> </u>	M00005491:812	CH02COH
1730	446514	RTA22200001F.a.16.1.P.Seq	F	M00042527:47	CH15CON
1731	456808	RTA22200010F.o.15.1.P.Seq	F	M00056512:59	CH16COP
1732	447035	RTA22200003F.f.08.1.P.Seq	F	M00055635:510	CH15CON
1733	446913	RTA22200001F.j.16.1.P.Seq	F	M00042742:16	CH15CON
1734	446900	RTA22200022F.g.20.1.P.Seq	F	M00054934:43	CH17COHLV
1735	504513	RTA22200003F.f.07.1.P.Seq	F	M00055635:810	CH15CON
1736	380477	RTA22200014F.j.09.1.P.Seq	F	M00056993:26	CH16COP
1737	1213	RTA22200244F.p.13.1.P.Seq	F	M00027231:13	CH04MAL
1738	8259	RTA22200222F.I.14.1.P.Seq	F	M00004093:310	CH01COH
1739	8259	RTA22200222F.h.23.1.P.Seq	F	M00003922:83	CH01COH
1740	552968	RTA22200019F.d.20.1.P.Seq	F	M00043503:38	CH17COHLV
1741	650845	RTA22200010F.o.09.1.P.Seq	F	M00056511:812	CH16COP
1742	648594	RTA22200014F.h.17.1.P.Seq	F	M00056977:73	CH16COP
1743	648594	RTA22200014F.h.17.2.P.Seq	F	M00056977:73	CH16COP
1744	2796	RTA22200240F.c.13.1.P.Seq	F	M00023331:111	CH04MAL
1745	5753	RTA22200227F.f.13.1.P.Seq	F	M00006648:15	CH02COH
1746	734256	RTA22200014F.I.11.2.P.Seq	F	M00057005:43	CH16COP
1747	449580	RTA22200007F.p.19.1.P.Seq	F	M00056290:58	CH15CON
1748	553705	RTA22200021F.a.23.2.P.Seq	F	M00054726:68	CH17COHLV
1749	730670	RTA22200011F.n.02.1.P.Seq	F	M00056638:21	CH16COP
1750	15035	RTA22200001F.c.07.1.P.Seq	F	M00042544:610	CH15CON
1751	394436	RTA22200002F.o.12.1.P.Seq	F	M00055552:39	CH15CON
1752	726810	RTA22200011F.h.22.1.P.Seq	F	M00056592:33	CH16COP
1753	352763	RTA22200022F.j.12.1.P.Seq	F	M00054949:17	CH17COHLV
1754	3506	RTA22200240F.i.14.1.P.Seq	F	M00023414:63	CH04MAL
1755	726377	RTA22200015F.i.14.1.P.Seq	F	M00057100:23	CH16COP
1756	562111	RTA22200018F.d.10.1.P.Seq	F	M00042460:17	CH17COHLV
1757	404475	RTA22200010F.m.04.1.P.Seq	F	M00056496:89	CH16COP
1758	13824	RTA22200234F.m.16.1.P.Seq	F	M00022370:72	CH03MAH
1759	558222	RTA22200019F.h.07.1.P.Seq	F	M00054502:41	CH17COHLV
1760	2834	RTA22200224F.p.09.1.P.Seq	F	M00005406:111	CH02COH
1761	453470	RTA22200001F.a.10.1.P.Seq	F	M00042523:35	CH15CON
1762	558682	RTA22200015F.o.16.1.P.Seq	F	M00057141:22	CH16COP
1763	641710	RTA22200016F.j.03.1.P.Seq	F	M00057231:79	CH16COP
1764	640221	RTA22200013F.k.12.1.P.Seq	F	M00056874:71	CH16COP
1765	559057	RTA22200016F.n.11.1.P.Seq	F	M00057270:89	CH16COP
1766	551433	RTA22200025F.m.12.2.P.Seq	F	M00055385:76	CH17COHLV
1767	5729	RTA22200228F.f.10.1.P.Seq	F	M00006819:25	CH02COH
1768	352763	RTA22200022F.g.09.1.P.Seq	F	M00054931:19	CH17COHLV
1769	375651	RTA22200012F.m.08.1.P.Seq	F	M00056757:63	CH16COP
1770	644032	RTA22200012F.m.04.1.P.Seq	F	M00056756:15	CH16COP
1771	185562	RTA22200240F.j.15.1.P.Seq	F	M00023428:611	CH04MAL
1772	736349	RTA22200014F.d.03.2.P.Seq	F	M00056947;69	CH16COP
1773	638870	RTA22200012F.h.13.1.P.Seq	F	M00056721:67	CH16COP
1774	649719	RTA22200012F.k.03.1.P.Seq	F	M00056739:48	CH16COP
1775	62016	RTA22200026F.d.09.1.P.Seq	F	M00055420:74	CH17COHLV
1776	2889	RTA22200020F.p.17.1.P.Seq	F	M000033420:74	CH02COH
1777	647135	RTA22200010F.k.08.1.P.Seq	F	M00007084.74	CH16COP
1778	8283	RTA222000101 .k.00.1.P.Seq	F	M00005309:55	CH02COH
1779	732121	RTA22200224F.g.10.1.F.Seq	F	M00005309:55 M00056949:66	CH16COP
1780	532307	RTA22200014F.u.00.1.F.Seq	F	M00054937:112	CH17COHLV
1780	6589	RTA22200022F.II.09.1.F.Seq	F	M00005810:67	CH02COH
1782	554678	RTA22200226F.II.24.1.P.Seq	F	M00056363:26	CH16COP
1702	204010	1717222000101.0.13.1.1.389	I -	พเบบบอบอบอ.20	CHIOCOP

Table 1

	ble 1				
SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	CONTRACTOR OF THE PROPERTY OF	LIBRARY
1783	450410	RTA22200021F.a.09.2.P.Seq	F	M00054722:41	CH17COHLV
1784	643924	RTA22200002F.a.21.1.P.Seq	F	M00055427:61	CH15CON
1785	453719	RTA22200002F.a.17.1.P.Seq	F	M00055426:86	CH15CON
1786	451811	RTA22200003F.k.17.1.P.Seq	F	M00055681:72	CH15CON
1787	453059	RTA22200003F.b.22.1.P.Seq	F	M00055586:62	CH15CON
1788	453457	RTA22200021F.a.06.2.P.Seq	F	M00054721:611	
1789	558454	RTA22200020F.a.20.1.P.Seq	F	M00054572:31	CH17COHLV
1790	417467	RTA22200025F.c.11.1.P.Seq	F	M00055289:42	CH17COHLV
1791	447850	RTA22200001F.a.23.1.P.Seq	F	M00042532:68	CH15CON
1792	557948	RTA22200024F.n.05.1.P.Seq	F	M00055245:19	CH17COHLV
1793	452685	RTA22200022F.b.14.1.P.Seq	F	M00054877:812	CH17COHLV
1794	446964	RTA22200019F.e.05.1.P.Seq	F	M00043504:78	CH17COHLV
1795	550318	RTA22200021F.d.04.2.P.Seq	F	M00054741:310	CH17COHLV
1796	407077	RTA22200023F.p.13.1.P.Seq	F	M00055134:23	CH17COHLV
1797	650864	RTA22200007F.o.19.1.P.Seq	F	M00056282:44	CH15CON
1798	644721	RTA22200003F.p.16.1.P.Seq	F	M00055731:812	-c
1799	485431	RTA22200013F.I.17.1.P.Seq	F	M00056885:36	CH16COP
1800	651073	RTA22200007F.f.05.1.P.Seq	F	M00056186:62	CH15CON
1801	725811	RTA22200012F.o.17.1.P.Seq	F	M00056776:49	CH16COP
1802	645139	RTA22200005F.k.02.1.P.Seg	F	M00055924:811	CH15CON
1803	185478	RTA22200248F.j.05.1.P.Seq	F	M00027578:65	CH04MAL
1804	1441	RTA22200228F.j.04.2.P.Seq	F	M00006859:44	CH02COH
1805	640005	RTA22200002F.i.16.1.P.Seq	F	M00055511:59	CH15CON
1806	728273	RTA22200015F.h.04.1.P.Seq	F	M00057091:34	CH16COP
1807	185579	RTA22200242F.e.07.1.P.Seq	F	M00026996:16	CH04MAL
1808	724473	RTA22200012F.f.23.1.P.Seq	F	M00056711:31	CH16COP
1809	559674	RTA22200014F.h.06.1.P.Seq	F	M00056974:64	CH16COP
1810	456026	RTA22200005F.d.13.1.P.Seq	F	M00055873:53	CH15CON
1811	549320	RTA22200022F.a.02.1.P.Seq	F	M00054867:37	CH17COHLV
1812	447338	RTA22200001F.k.04.1.P.Seq	F	M00042746:29	CH15CON
1813	560700	RTA22200002F.b.09.1.P.Seq	F	M00055430:82	CH15CON
1814	3070	RTA22200021F.g.19.3.P.Seq	F	M00054769:67	CH17COHLV
1815	3070	RTA22200002F.f.03.1.P.Seq	F	M00055463:810	CH15CON
1816	380477	RTA22200014F.j.09.2.P.Seq	F	M00056993:26	CH16COP
1817	735040	RTA22200015F.b.05.1.P.Seq	F F	M00057047:32	CH16COP
1818	378525	RTA22200009F.j.16.2.P.Seq	F	M00042826:33	CH16COP
1819	284586	RTA22200009F.c.23.2.P.Seq	F	M00042756:21	CH16COP
1820	640276	RTA22200014F.e.15.2.P.Seq	F.	M00056956:61	CH16COP
1821	3344	RTA22200230F.j.07.1.P.Seq	F	M00007178:43	CH02COH
1822	555830	RTA22200005F.b.23.1.P.Seq	F	M00055861:43	CH15CON
1823	726307	RTA22200011F.i.10.1.P.Seq	F	M00056595:62	CH16COP
1824	416	RTA22200011F.m.06.1.P.Seq	F	M00056631:75	CH16COP
1825	2543	RTA22200225F.o.10.1.P.Seq	F	M00000051:73	CH02COH
1826	639352	RTA22200001F.n.09.1.P.Seq	F	M00042908:71	CH15CON
1827	453592	RTA222000011.11.03.11.F.Seq	F	M00042842:31	CH16COP
1828	450633	RTA22200009F.n.19.2.P.Seq	F	M00042842.51	CH15CON
1829	448383	RTA22200000F.II.15.2.P.Seq	F	M00038107.38	CH16COP
1830	648719	RTA22200008F.a.08.1.P.Seq	F	M00056293:65	CH16COP CH15CON
1831	730655	RTA22200008F.a.08.1.F.Seq	F	M00056722:71	CH16COP
1832	141185	RTA22200012F.II.10.1.P.Seq	F.	M00056228:17	CH15CON
1833	640498	RTA22200007F.J.21.2.P.Seq	F [		
1834	9029	RTA22200003F.III.07.1.P.Seq	F	M00055703:28	CH15CON
1835	559674	RTA22200222F.1.06.1.P.Seq	F i	M00004087:211	CH01COH
1836	555734	RTA22200014F.n.09.2.P.Seq	F	M00056974:64	CH16COP
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Table 1

	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
1837	1943	RTA22200017F.d.22.1.P.Seq	F	M00057333:69	CH16COP
1838	648320	RTA222000171.d.22.111.3eq	<u> </u>	M00057353.33	CH15CON
1839	558098	RTA22200024F.p.10.1.P.Seq	F	M00055260:15	CH17COHLV
1840	468672	RTA2220004F.h.23.1.P.Seq	F	M00055794:26	CH15CON
1841	456596	RTA22200022F.f.19.1.P.Seq	F F	M00054926:61	CH17COHLV
1842	649722	RTA22200012F.n.07.1.P.Seq	F	M00054520:01	CH16COP
1843	550708	RTA22200012F.0.19.1.P.Seq	F	M00054996:211	Secretaria de la constante de
1844	643931	RTA22200022F.0.19.1.F.Seq	F	M00054990.211	CH15CON
1845	726927	RTA22200031 II.14-1.1 .3.64	F	M00056603:41	CH16COP
1846	459012	RTA2220007F.b.13.1.P.Seq	F	M00056005.41	CH15CON
1847	397773	RTA22200007 .5.13.1.P.Seq	F	M00055592:15	CH15CON
1848	450004	RTA222000011F.f.04.1.P.Seq	F	M00056566:81	CH16COP
1849	649732	RTA2220007F.o.23.1.P.Seq	F	M00056380:81	CH15CON
1850	553955	RTA222000071.0.23.1.F.3eq		M00050202:09	CH17COHLV
1851	646309	RTA222000217.1.08.3.F.Seq	F	M00054755:86 M00056718:21	CH16COP
1852	**************************************	RTA22200012F.II.04.1.F.Seq	F	M00055718.21	CH17COHLV
1853	402727	RTA22200024F.m.17.1.P.Seq	F	M00055243:71	CH17COHLV
Incompany mount on a company	468736	RTA22200024F.III.17.1.P.Seq	F	M00055243.71	CH17CORLV
1854	650422	RTA22200007F.a.16.1.P.Seq	F		&
1855	730533 726307	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	F	M00056808:212	CH16COP
1856		RTA22200015F.a.17.1.P.Seq	F	M00057044:66	CH16COP
1857	450311	RTA22200005F.j.22.1.P.Seq	F	M00055922:32	CH15CON
1858	450940	RTA22200001F.j.15.1.P.Seq	F	M00042742:83	CH15CON
1859	726786	RTA22200015F.f.06.1.P.Seq		M00057077:42	CH16COP
1860	7634	RTA22200235F.b.19.1.P.Seq	<u> </u>	M00022420:83	CH03MAH
1861	230995	RTA22200016F.e.14.1.P.Seq	<u> </u>	M00057196:64	CH16COP
1862	374770	RTA22200003F.n.21.1.P.Seq	<u> </u>	M00055720:16	CH15CON
1863	9275	RTA22200249F.g.11.1.P.Seq	E	M00027665:41	CH04MAL
1864	553860	RTA22200023F.k.02.1.P.Seq	F F	M00055077:611	<u></u>
1865	452010	RTA22200001F.m.07.1.P.Seq	Same and the second	M00042900:37	CH15CON
1866	649560	RTA22200015F.j.16.1.P.Seq	F	M00057106:52	CH16COP
1867	452704	RTA22200008F.b.18.1.P.Seq	F F	M00056307:812	CH15CON
1868	447594	RTA22200007F.j.10.1.P.Seq	F	M00056225:79 M00057130:811	CH15CON
1869 1870	555444	RTA22200015F.n.10.1.P.Seq	F	M00057318:13	CH16COP
	736556	RTA22200017F.c.20.1.P.Seq	F		CH16COP
1871	5289	RTA22200235F.I.12.2.P.Seq	F	M00022533:56	CH03MAH
1872	732121	RTA22200015F.c.11.1.P.Seq	F	M00057059:66	CH16COP
1873	452567	RTA22200009F.f.12.1.P.Seq	F	M00042783:13	CH16COP
1874	551634	RTA22200002F.n.04.1.P.Seq		M00055544:54	CH15CON
1875	644099	RTA22200005F.g.07.1.P.Seq	F.	M00055887:67	CH15CON
1876	726788	RTA22200012F.h.08.1.P.Seq	<u> </u>	M00056719:19	CH16COP
1877	638802	RTA22200012F.a.02.1.P.Seq	<u> </u>	M00056662:63	CH16COP
1878	646283	RTA22200002F.o.18.1.P.Seq	<u> </u>	M00055553:82	CH15CON
1879	8403	RTA22200238F.n.03.1.P.Seq	F	M00022998:38	CH03MAH
1880	2224	RTA22200233F.g.23.1.P.Seq	Ę	M00008079:33	CH03MAH
1881	650053	RTA22200008F.g.02.1.P.Seq	F	M00056459:47	CH15CON
1882	380477	RTA22200008F.e.20.1.P.Seq	<u>F</u>	M00056437:87	CH15CON
1883	450867	RTA22200019F.a.09.1.P.Seq	<u> </u>	M00043386:16	CH17COHLV
1884	456764	RTA22200003F.n.22.1.P.Seq	F -	M00055720:11	CH15CON
1885	641373	RTA22200003F.b.11.1.P.Seq	F	M00055583:24	CH15CON
1886	555882	RTA22200004F.m.12.1.P.Seq	<u> </u>	M00055818:510	CH15CON
1887	644046	RTA22200008F.b.04.1.P.Seq	F	M00056302:612	CH15CON
1888	447250	RTA22200009F.d.15.2.P.Seq	F	M00042760:37	CH16COP
1889	456596	RTA22200022F.o.14.1.P.Seq	F	M00054995:310	CH17COHLV
1890	2218	RTA22200225F.d.15.1.P.Seq	F	M00005447:41	CH02COH

Table 1

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and the second second second second	CLUSTER	SEQ NAME	ORIENTATION	AND A STREET WAS A	LIBRARY
1891	446450	RTA22200001F.h.16.1.P.Seq	F	M00042718:33	CH15CON
1892	640889	RTA22200003F.I.12.1.P.Seq	F	M00055691:57	CH15CON
1893	530774	RTA22200025F.n.12.2.P.Seq	F	M00055391:78	CH17COHLV
1894	649062	RTA22200002F.j.14.1.P.Seq	F	M00055519:87	CH15CON
1895	12808	RTA22200233F.m.15.1.P.Seq	F	M00021649:512	CH03MAH
1896	468672	RTA22200001F.g.11.1.P.Seq	F	M00042704:64	CH15CON
1897	650773	RTA22200008F.d.12.1.P.Seq	F	M00056324:42	CH15CON
1898	732237	RTA22200014F.j.12.2.P.Seq	F	M00056994:67	CH16COP
1899	650773	RTA22200001F.o.15.1.P.Seq	F	M00054798:61	CH15CON
1900	550216	RTA22200021F.b.09.2.P.Seq	F	M00054727:86	CH17COHLV
1901	639189	RTA22200002F.n.20.1.P.Seq	F	M00055547:55	CH15CON
1902	3447	RTA22200227F.I.17.1.P.Seq	F	M00006719:87	CH02COH
1903	2012	RTA22200248F.c.03.1.P.Seq	F	M00027524:211	CH04MAL
1904	642876	RTA22200005F.b.13.1.P.Seq	F	M00055856:64	CH15CON
1905	449690	RTA22200009F.j.10.2.P.Seq	F	M00042823:74	CH16COP
1906	451208	RTA22200004F.k.07.1.P.Seq	F	M00055804:63	CH15CON
1907	725811	RTA22200011F.k.18.1.P.Seq	F	M00056616:38	CH16COP
1908	1256	RTA22200232F.b.11.1.P.Seq	F	M00021869:41	CH03MAH
1909	446599	RTA22200004F.f.18.1.P.Seq	F	M00055780:78	CH15CON
1910	446537	RTA22200001F.c.05.1.P.Seq	F	M00042544:42	CH15CON
1911	726281	RTA22200010F.l.02.1.P.Seq	F	M00056484:72	CH16COP
1912	11286	RTA22200226F.n.17.1.P.Seq	F	M00005802:810	CH02COH
1913	556082	RTA22200022F.p.11.1.P.Seq	F	M00055005:811	CH17COHLV
1914	97507	RTA22200005F.a.21.1.P.Seq	F	M00055851:711	CH15CON
1915	535955	RTA22200022F.b.11.1.P.Seq	F	M00054876:73	CH17COHLV
1916	728251	RTA22200011F.i.08.1.P.Seq	F	M00056595:37	CH16COP
1917	733849	RTA22200015F.g.08.1.P.Seq	F	M00057085:13	CH16COP
1918	447574	RTA22200014F.k.20.2.P.Seq	F	M00057002:25	CH16COP
1919	7607	RTA22200229F.g.17.1.P.Seq	F	M00006976:45	CH02COH
1920	644032	RTA22200010F.i.21.1.P.Seq	F	M00056425:23	CH16COP
1921	454087	RTA22200012F.f.13.1.P.Seq	F	M00056709:15	CH16COP
1922	412364	RTA22200007F.p.02.1.P.Seq	F	M00056283:33	CH15CON
1923	535208	RTA22200002F.m.04.1.P.Seq	F	M00055534:81	CH15CON
1924	644609	RTA22200002F.m.21.1.P.Seq	F	M00055542:66	CH15CON
1925	645073	RTA22200004F.o.14.1.P.Seq	F	M00055830:710	CH15CON
1926	417467	RTA22200012F.g.08.1.P.Seq	F	M00056712:26	CH16COP
1927	554188	RTA22200004F.n.02.1.P.Seq	F	M00055821:712	CH15CON
1928	647185	RTA22200005F.n.02.1.P.Seq	F	M00055958:62	CH15CON
1929	736679	RTA22200012F.a.20.1.P.Seq	F	M00056666:38	CH16COP
1930	553547	RTA22200022F.j.02.1.P.Seq	F	M00054947:712	CH17COHLV
1931	641524	RTA22200016F.c.06.1.P.Seq	F	M00057170:83	CH16COP
1932	649717	RTA22200003F.f.02.1.P.Seq	F	M00055633:12	CH15CON
1933	451041	RTA22200018F.a.16.1.P.Seq	F	M00042355:31	CH17COHLV
1934	3483	RTA22200225F.e.24.1.P.Seq	F	M000042050:31	CH02COH
1935	500959	RTA2220008F.c.24.1.P.Seq	F	M00056323:810	CH15CON
1936	500959	RTA22200008F.d.01.1.P.Seq	F	M00056323:810	CH15CON
1937	697	RTA2220003F.d.01.1.F.Seq	F	M000030323.810	CH03MAH
1938	736955	RTA22200233F.f.22.1.P.Seq	F	M00056840:89	CH16COP
1938	554742	RTA22200013F.1.22.1.F.Seq	F	M00055811:312	CH15CON
1939	642973	RTA22200004F.I.14.1.P.Seq	F	M00057132:68	CH16COP
1940	449437	RTA22200013F.h.14.1.P.Seq	F	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
1941	449437 467991		F F	M00056304:711	CH15CON
1942	650204	RTA22200002F.j.11.1.P.Seq RTA22200004F.n.13.1.P.Seq	F F	M00055517:43	CH15CON
1943	640618	RTA22200004F.n.13.1.P.Seq	F	M00055825:53	CH15CON
1344	040010	KIAZZZUUUU4F.p.17.1.P.Seq		M00055840:46	CH15CON

Table 1

10	ble 1				
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1946	640276	RTA22200012F.p.05.1.P.Seq	F	M00056780:69	CH16COP
1947	554101	RTA22200012F.f.24.1.P.Seq	F	M00056711:65	CH16COP
1948	185432	RTA22200249F.p.01.1.P.Seq	F	M00027742:21	CH04MAL
1949	455598	RTA22200022F.m.06.1.P.Seq	F	M00054970:73	CH17COHLV
1950	649354	RTA22200001F.o.05.1.P.Seq	F	M00054792:59	CH15CON
1951	4408	RTA22200242F.j.13.1.P.Seq	F	M00027034:49	CH04MAL
1952	452366	RTA22200006F.o.03.2.P.Seq	F	M00056110:49	CH15CON
1953	452366	RTA22200001F.e.02.1.P.Seq	F	M00042563:79	CH15CON
1954	727331	RTA22200016F.a.19.1.P.Seq	F	M00057161:59	CH16COP
1955	644853	RTA22200008F.a.05.1.P.Seq	F	M00056293:58	CH15CON
1956	554079	RTA22200014F.k.21.2.P.Seq	F	M00057002:26	CH16COP
1957	556245	RTA22200023F.I.05.1.P.Seq	F	M00055088:112	CH17COHLV
1958	557388	RTA22200012F.p.13.1.P.Seq	F	M00056783:711	CH16COP
1959	449468	RTA22200012F.I.11.1.P.Seq	F	M00056752:51	CH16COP
1960	556245	RTA22200023F.o.12.1.P.Seq	F	M00055125:56	CH17COHLV
1961	455327	RTA22200013F.b.03.1.P.Seq	F	M00056804:56	CH16COP
1962	546632	RTA22200015F.a.10.1.P.Seq	F	M00057041:211	CH16COP
1963	558762	RTA22200022F.a.03.1.P.Seq	F	M00054867:22	CH17COHLV
1964	550818	RTA22200020F.d.10.1.P.Seq	F	M00054590:72	CH17COHLV
1965	554079	RTA22200021F.p.09.2.P.Seq	F	M00054862:27	CH17COHLV
1966	452430	RTA22200016F.e.07.1.P.Seq	F	M00057192:52	CH16COP
1967	452430	RTA22200008F.e.11.1.P.Seq	F F	M00056342:611	CH15CON
1968	556082	RTA22200019F.i.02.1.P.Seq	F	M00054507:311	CH17COHLV
1969	514418	RTA22200023F.e.23.1.P.Seq	F	M00055046:511	CH17COHLV
1970	426895	RTA22200021F.p.14.2.P.Seq	F	M00054863:73	CH17COHLV
1971	560803	RTA22200025F.c.05.1.P.Seq	F	M00055283:65	CH17COHLV
1972	447737	RTA22200018F.c.11.1.P.Seq	F	M00042453:51	CH17COHLV
1973	373432	RTA22200018F.I.18.1.P.Seq	F	M00042400:01	CH17COHLV
1974	779	RTA22200250F.f.02.1.P.Seq	F	M00027820:32	CH04MAL
1975	455327	RTA22200015F.m.05.1.P.Seg	F	M00057124:410	CH16COP
1976	554742	RTA22200004F.k.16.1.P.Seq	F	M00055806:812	CH15CON
1977	455327	RTA22200022F.g.14.1.P.Seq	F	M00054933:47	CH17COHLV
1978	11043	RTA22200228F.j.10.2.P.Seq	F	M00004000:47	CH02COH
1979	727447	RTA22200012F.c.14.1.P.Seq	F	M00056684:15	CH16COP
1980	552905	RTA22200011F.i.16.1.P.Seq	F F	M00056597:67	CH16COP
1981	446900	RTA22200001F.g.23.1.P.Seq	F	M00042711:711	CH15CON
1982	644190	RTA22200005F.j.02.1.P.Seq	F	M00055913:45	CH15CON
1983	455327	RTA22200021F.c.18.2.P.Seq	F	M00054739:73	CH17COHLV
1984	422375	RTA22200020F.g.01.1.P.Seq	F	M00054613:49	CH17COHLV
1985	422375	RTA22200020F.f.24.1.P.Seq	F	M00054613:49	CH17COHLV
1986	530774	RTA22200025F.p.22.2.P.Seq	F	M00055405:49	CH17COHLV
1987	554101	RTA222000231 .p.22.2.F .Seq	F	M00055405.49	CH16COP
1988	5268	RTA222000121.g.0111.F.Seq	F	M000056711:05	CH02COH
1989	642461	RTA22200220F.g.10.1.F.Seq	F	M000056715:71	CH16COP
1990	770	RTA22200012F.g.16.1.F.Seq	F	M00055575:48	CH15CON
1990	3837	RTA22200003F.a.16.1.F.Seq	F	M00003373.48	CH03MAH
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1992	561382 4408	RTA22200022F.d.20.1.P.Seq	F	M00054899:71 M00027543:29	CH17COHLV CH04MAL
1993	4408	RTA22200248F.e.13.1.P.Seq	F	**************************************	
1994	5686	RTA22200234F.j.21.1.P.Seq	·	M00022279:84	CH03MAH
1995	374609	RTA22200013F.e.23.1.P.Seq	<u> </u>	M00056833:31	CH16COP
1996	734793	RTA22200012F.h.06.1.P.Seq	<u> </u>	M00056719:46	CH16COP
1997	452430	RTA22200014F.f.18.2.P.Seq	F	M00056964:12	CH16COP
1998	450940	RTA22200020F.n.11.1.P.Seq	F	M00054684:812	CH17COHLV

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CONTRACTOR	CLUSTER	SEQ NAME	ORIENTATION	complete announcement for a service of the contract of the con	LIBRARY
1999	460445	RTA22200024F.c.23.1.P.Seq	F	M00055162:66	CH17COHLV
2000	549041	RTA22200020F.i.01.1.P.Seq	F	M00054636:23	CH17COHLV
2001	555276	RTA22200018F.g.13.1.P.Seq	F	M00043313:77	CH17COHLV
2002	426895	RTA22200015F.I.18.1.P.Seq	- F	M00057120:512	CH16COP
2003	1833	RTA22200224F.f.16.1.P.Seq	F	M00005293:46	CH02COH
2004	446450	RTA22200009F.h.08.1.P.Seq	F	M00042805:56	CH16COP
2005	650517	RTA22200007F.I.18.2.P.Seq	F	M00056248:19	CH15CON
2006	554785	RTA22200026F.f.14.1.P.Seq	F	M00055479:312	karananan maranan mara
2007	607430	RTA22200002F.i.06.1.P.Seq	F	M00055508:53	CH15CON
2008	446673	RTA22200012F.a.18.1.P.Seq	F	M00056665:55	CH16COP
2009	734685	RTA22200014F.i.17.2.P.Seq	F	M00056986:71	CH16COP
2010	11630	RTA22200248F.h.02.1.P.Seq	F	M00027562:32	CH04MAL
2011	2930	RTA22200228F.c.13.1.P.Seq	F	M00006795:47	CH02COH
2012	44424	RTA22200006F.f.07.2.P.Seq	F	M00056038:64	CH15CON
2013	452052	RTA22200001F.e.19.1.P.Seq	F	M00042574:88	CH15CON
2014	449356	RTA22200009F.j.14.2.P.Seq	F	M00042826:35	CH16COP
2015	726225	RTA22200010F.d.04.1.P.Seq	F	M00056372:310	CH16COP
2016	453708	RTA22200022F.j.14.1.P.Seq	F	M00054950:76	CH17COHLV
2017	447858	RTA22200003F.n.20.1.P.Seq	F	M00055720:79	CH15CON
2018	451613	RTA22200018F.e.18.1.P.Seq	F	M00043301:89	CH17COHLV
2019	650337	RTA22200008F.c.18.1.P.Seq	F	M00056319:71	CH15CON
2020	62016	RTA22200019F.k.10.1.P.Seq	F	M00054522:811	CH17COHLV
2021	447250	RTA22200014F.f.02.2.P.Seq	F	M00056960:35	CH16COP
2022	3837	RTA22200231F.c.01.1.P.Seq	F	M00007936:29	CH03MAH
2023	640614	RTA22200017F.e.22.1.P.Seg	F	M00057340:312	CH16COP
2024	729531	RTA22200013F.g.24.1.P.Seq	F	M00056850:511	CH16COP
2025	729531	RTA22200013F.h.01.1.P.Seq	F	M00056850:511	CH16COP
2026	647952	RTA22200004F.j.07.1.P.Seq	F	M00055800:38	CH15CON
2027	446913	RTA22200004F.k.19.1.P.Seq	F	M00055806:67	CH15CON
2028	2675	RTA22200233F.a.20.1.P.Seq	F	M00008015:68	CH03MAH
2029	643481	RTA22200003F.i.13.1.P.Seq	F	M00055664:18	CH15CON
2030	1345	RTA22200012F.I.22.1.P.Seq	F	M00056754:84	CH16COP
2031	26	RTA22200231F.a.03.1.P.Seq	F	M00007926:15	CH03MAH
2032	945	RTA22200230F.h.02.1.P.Seq	F	M00007156:56	CH02COH
2033	449169	RTA22200009F.b.09.2.P.Seq	F	M00042461:110	CH16COP
2034	394193	RTA22200007F.h.10.1.P.Seq	F	M00056209:612	CH15CON
2035	452212	RTA22200006F.h.13.2.P.Seq	F	M00056055:21	CH15CON
2036	394193	RTA22200007F.i.15.2.P.Seq	F	M00056220:42	CH15CON
2037	1310	RTA22200235F.o.10.2.P.Seq	F	M00022561:16	CH03MAH
2038	734094	RTA22200016F.k.07.1.P.Seq	F	M00057241:24	CH16COP
2039	646579	RTA22200002F.h.08.1.P.Seq	F	M00055498:89	CH15CON
2040	4471	RTA22200233F.j.22.1.P.Seq	F	M00008098:81	CH03MAH
2041	729173	RTA22200016F.e.13.1.P.Seq	F	M00057196:53	CH16COP
2042	450323	RTA22200009F.i.11.2.P.Seq	F	M00042814:211	CH16COP
2043	4652	RTA22200241F.n.17.1.P.Seq	F	M00026950:612	CH04MAL
2044	553316	RTA22200021F.a.10.2.P.Seq	F	M00054722:38	CH17COHLV
2045	642604	RTA22200016F.j.10.1.P.Seq	F	M00057233:67	CH16COP
2046	553316	RTA22200002F.i.02.1.P.Seq	F	M00055504:48	CH15CON
2047	4097	RTA22200249F.f.14.1.P.Seq	F	M00027652:73	CH04MAL
2048	6818	RTA22200006F.f.12.2.P.Seq	F	M00056042:11	CH15CON
2049	395341	RTA22200024F.c.22.1.P.Seq	F F	M00055161:111	CH17COHLV
2050	649143	RTA22200007F.o.20.1.P.Seq	F I	M00056282:31	CH15CON
2051	649143	RTA22200007F.o.18.1.P.Seq	F	M00056282:611	CH15CON
2052	648310	RTA22200007F.e.16.1.P.Seq	F.	M00056184:61	CH15CON
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Table 1

	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
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2055	6878	RTA22200004F.B.12.1.F.Seq	F	M00003744.08	CH03MAH
2056	452238	RTA222002511.d.0111.7.3eq	F	M00007320.17 M00042758:812	CH16COP
2057	1870	RTA22200238F.g.09.1.P.Seq	F	M00042730.012	CH03MAH
2058	559259	RTA22200002F.n.12.1.P.Seq	F.	M00055545:812	CH15CON
2059	453457	RTA22200004F.a.06.1.P.Seq	F	M00055735:37	CH15CON
2060	8868	RTA22200033F.I.13.1.P.Seq	F	M00033733:37 M00021626:64	CH03MAH
2061	453059	RTA22200001F.b.21.1.P.Seq	F	M00021020:04 M00042540:63	CH15CON
2062	236368	RTA222000011.5.21111.5eq	F	M00042340:03 M00056323:17	CH15CON
2063	453059	RTA22200001 .d.03.11 .Geq	F	M00030323:17	CH15CON
2064	549979	RTA22200007F.c.06.1.P.Seq	F	M00042353:35 M00056153:41	CH15CON
2065	515631	RTA22200011 .c.00.1.F .Seq	F	M00056996:112	CH16COP
2066	2235	RTA222000141 J.22.2.1 .Seq	F	M00030330:112	CH03MAH
2067	448193	RTA22200234F.1.22.1.F.Seq	F	M00057271:51	CH16COP
2068	530774	RTA22200010F.c.12.1.P.Seq	F	M00057271:51	CH16COP
2069	650204	RTA22200010F.C.12.1.F.Seq	F	M00055450:28	CH15CON
2070	644240	RTA22200005F.j.19.1.P.Seq		M00055430.28	CH15CON
2071	552614	RTA222000031 J. 19.1.F. Seq	F	M00053921:35 M00054965:82	CH17COHLV
2072	727331	RTA22200022F.I.TT.I.F.Seq	F	M00054905.82 M00056532:76	CH16COP
2072	185457	RTA22200011F.b.04.1.F.Seq	F	M00030332.76	CH04MAL
2073	454531	RTA22200244F.II.03.1.F.Seq	F	M00054869:41	CH17COHLV
2075	643485	RTA22200022F.a.11.1.F.Seq	F	M00054869.41	CH15CON
2075	733669	RTA222000013F.m.22.1.P.Seq	F	M00056895:17	CH16COP
2077	452344	RTA22200013F.III.22.1.F.Seq	F	M00050895.17	CH16COP
2078	63602	RTA22200010F.b.19.1.F.Seq	F	M00057167.27	CH15CON
2079	454155	RTA22200002F.J.11.1.F.Seq	F	M00055802:48	CH15CON
2019	549903	RTA22200004F.j.13.1.F.Seq	F	M00055802.48	CH16COP
2081	515631	RTA22200013F.b.22.1.F.3eq	F	M00056996:112	CH16COP
2082	6878	RTA222000141 j.22.1.1 .Seq	F	M000030390:112	CH04MAL
2083	2977	RTA22200231F.I.18.1.P.Seq	F	M000027300:310	CH03MAH
2084	553823	RTA22200006F.i.02.2.P.Seq	F	M00056060:210	CH15CON
2085	3070	RTA22200026F.a.07.1.P.Seq	F	M00055405:77	CH17COHLV
2086	728884	RTA22200012F.k.06.1.P.Seq	F	M00055405:77	CH16COP
2087	8166	RTA22200009F.c.03.2.P.Seq	F	M00030740:23	CH16COP
2088	644190	RTA22200010F.b.18.1.P.Seq	F	M00056362:75	CH16COP
2089	733669	RTA22200015F.I.05.1.P.Seq	F	M00057117:711	CH16COP
2090	728273	RTA22200011F.i.07.1.P.Seq	F	M00056595:12	MOREO PARAMENTA POR MISARCINA MARTINA RAMANA MARTINA PARAMENTA ANTONIO PARAMENTA PARAM
2091	406499	RTA22200004F.a.13.1.P.Seq	F	M00055736:77	CH15CON
2092	557720	RTA22200022F.k.07.1.P.Seq	F	M00054954:61	CH17COHLV
2093	732050	RTA22200015F.d.10.1.P.Seq	F	M00057065:44	CH16COP
2094	450867	RTA222000131.d.10.11.F.Seq	F	M00057663.44 M00054648:810	CH17COHLV
2095	650297	RTA222000201 j. 10. 1. F. Seq	F	M00054048.810	CH16COP
2096	448064	RTA222000101 .a.01.1.F.Seq	F	M00037131.24 M00042781:16	CH16COP
2097	452530	RTA222000031.1.00.1.10eq	F	M00042701:10	CH17COHLV
2098	7592	RTA222000101 .p.03.1.F.Seq	F	M00045574.20	CH02COH
2099	733669	RTA22200225F.III.10.1.F.Seq	F	M000057165:512	CH16COP
2100	11028	RTA222000101.b.13.1.F.Seq	F	M000037103.312	CH02COH
2101	1013	RTA22200001F.h.22.1.P.Seq	F	M00042720:710	CH15CON
2102	549265	RTA222000017.11.22.1.17.Seq	F	M00042720.710	CH15CON
2102	376600	RTA22200002F.II.03.1.F.Seq	F	M00056911:62	CH16COP
2103	643804	RTA222000131.p.0d.1.F.Seq	F	M00055839:110	CH15CON
2105	454927	RTA22200004F.p.14.1.F.3eq	F	M00055839.110	CH17COHLV
2106	446528	RTA22200026F.i.06.2.P.Seq	F	M00055420.610	CH15CON
2100		1717222000001 1,00.2.F .3eq		17100000001100	OFFICEN

Table 1

SEQ ID   CLUSTER   SEQ NAME   ORIENTATION   CLONE ID   LIBRARY   2107   2218   RTA22200026E n.14.1.P.Seq   F   M0005590.411   CH02COH		CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
2108					THE SECRET PLANS AND ADDRESS OF THE PARTY OF	J
2110 84895 RTA22200014F.m13.2.P.Seq F M00057013.41 CH16COP 2111 19509 RTA22200225F.h22.1.P.Seq F M00006890.52 CH02COH 2112 7037 RTA22200230F.k.08.1.P.Seq F M00006890.52 CH02COH 2113 559906 RTA22200025F.h22.1.P.Seq F M000057194.24 CH02COH 2114 452076 RTA22200004F.g.03.1.P.Seq F M00055705.0110 CH16CON 2114 452076 RTA22200004F.g.03.1.P.Seq F M00055705.0110 CH16CON 2114 452076 RTA22200004F.g.03.1.P.Seq F M00055783.311 CH15CON 2116 559974 RTA22200004F.g.03.1.P.Seq F M00055771.44 CH15CON 2117 2235 RTA22200232F.n.21.1.P.Seq F M00055771.12 CH17COHLV 2118 7545 RTA22200023F.n.21.1.P.Seq F M0005771.12 CH17COHLV 2119 729173 RTA222000216F.n.22.1.P.Seq F M0005777.12 CH17COHLV 2120 650448 RTA22200004F.m.09.1.P.Seq F M0005577.11 CH16COP 2121 172013 RTA22200004F.m.09.1.P.Seq F M00055818.612 CH15CON 2122 651088 RTA22200014F.c.15.1.P.Seq F M00056845.211 CH16COP 2123 651088 RTA22200014F.c.15.1.P.Seq F M00056845.211 CH16COP 2124 728810 RTA22200014F.c.15.1.P.Seq F M00056845.211 CH16COP 2125 406499 RTA22200014F.c.15.1.P.Seq F M00056845.211 CH16COP 2126 556326 RTA22200004F.n.03.1.P.Seq F M000568945.211 CH16COP 2127 644836 RTA22200004F.n.03.1.P.Seq F M000568945.211 CH16COP 2128 649062 RTA22200004F.n.03.1.P.Seq F M000568945.211 CH16COP 2128 649062 RTA22200001F.0.06.1.P.Seq F M00056897.11 CH17COHLV 2131 72813 RTA22200001F.0.06.1.P.Seq F M00056897.11 CH17COHLV 2131 72813 RTA2220001F.0.06.1.P.Seq F M00056897.11 CH17COHLV 2131 72813 RTA2220001F.0.06.1.P.Seq F M0005697.11 CH17COHLV 2131 72813 RTA2220001F.0.08.1.P.Seq F M0005690.11 CH16COP 2133 727314 RTA2220001F.0.08.2.P.Seq F M0005690.11 CH16COP 2134 65603 RTA2220001F.0.08.2.P.Seq F M0005690.11 CH16COP 2135 651382 RTA2220001F.0.08.2.P.Seq F M0005690.11 CH16COP 2136 646883 RTA2220001F.0.08.P.Seq F M0005690.11 CH16COP 2137 644868 RTA2220001F.0.08.P.Seq F M0	Access was a construction of the construction	in and the second secon	reactive contract and the second seco		The her wind his home will have a committee a committee to be a co	de reconstruire de la company
2110 157629 RTA22200242F.c.11.1.P.Seq F M0006990:52 CH02COH 2112 2930 RTA22200239F.h.08.1.P.Seq F M00007194:24 CH02COH 2113 559806 RTA22200002F.i04.1.P.Seq F M00005505:110 CH15CON 2114 452076 RTA22200002F.j.04.1.P.Seq F M00055797:44 CH15CON 2115 454869 RTA22200004F.e.14.1.P.Seq F M00055771:44 CH15CON 2116 559674 RTA22200023F.j.05.1.P.Seq F M00055771:44 CH15CON 2116 559674 RTA22200023F.j.05.1.P.Seq F M00055771:44 CH15CON 2117 2235 RTA2220023F.n.21.1.P.Seq F M00055771:42 CH17COHLV 2117 72235 RTA2220023F.n.21.1.P.Seq F M00055771:42 CH17COHLV 2119 729173 RTA22200023F.n.22.1.P.Seq F M0002494:31 CH16COP 2120 650448 RTA22200024F.m.09.1.P.Seq F M00055772:11 CH16COP 2120 650448 RTA22200014F.c.15.1.P.Seq F M00055727:11 CH16COP 2120 650488 RTA22200014F.c.15.1.P.Seq F M00056945:211 CH16COP 2123 651088 RTA22200014F.c.15.1.P.Seq F M00056945:211 CH16COP 2124 728810 RTA22200014F.c.15.2.P.Seq F M00056945:211 CH16COP 2126 556325 RTA22200004F.n.23.1.P.Seq F M00056945:211 CH16COP 2126 464836 RTA22200004F.n.23.1.P.Seq F M00056945:211 CH16COP 2126 464836 RTA22200004F.n.23.1.P.Seq F M00056977:48 CH15CON 2129 454776 RTA22200004F.n.23.1.P.Seq F M00055977:48 CH15CON 2129 454776 RTA22200004F.n.23.1.P.Seq F M00056937:110 CH17COHLV 2127 644836 RTA22200004F.n.23.1.P.Seq F M00055977:48 CH15CON 2129 454776 RTA22200014F.o.95.1.P.Seq F M00055977:48 CH15CON 2129 454776 RTA22200014F.o.95.1.P.Seq F M0005693:2:71 CH17COHLV 2127 744848 RTA22200014F.n.23.1.P.Seq F M0005695:92 CH15CON 2129 454776 RTA22200014F.o.95.1.P.Seq F M0005693:2:71 CH17COHLV 2127 74484 RTA22200014F.o.95.1.P.Seq F M0005695:92 CH15CON 2133 77314 RTA22200014F.o.95.1.P.Seq F M00057024:75 CH16COP 2136 735579 RTA22200016F.a.05.1.P.Seq F M00057024:75 CH16COP 2136 735579 RTA22200016F.a.05.1.P.Seq F M00057024:75 CH16COP 2136 735579 RTA22200016F.a.05.1.P.Seq F M0005693:2:10 CH16COP 2144 40368 RTA22200014F.	\$				} {************************************	\$
2111   2930   RTA22200229F h.22.1 P.Seq	\$	·				· .
2112 7037 RTA22200230F.K08.1P.Seq F M00055505:110 CH15CON 2114 452076 RTA22200002F.Io.41.P.Seq F M00055783:311 CH15CON 2115 454869 RTA22200004F.e.14.1P.Seq F M00055783:311 CH15CON 2116 454869 RTA2220003F.j.05.1P.Seq F M0005571:44 CH15CON 2116 559674 RTA22200023F.j.05.1P.Seq F M0005771:42 CH17COHLV 2117 2235 RTA2220023F.j.05.1P.Seq F M0005771:41 CH15CON 2118 7545 RTA22200222F.k.06.1P.Seq F M0005071:12 CH17COHLV 2119 729173 RTA22200022F.k.06.1P.Seq F M00050571:2 CH17COHLV 2119 729173 RTA2220004F.m.29.1P.Seq F M00056818:612 CH15CON 2120 656448 RTA22200024F.m.29.1P.Seq F M00056818:612 CH15CON 2121 172013 RTA22200016F.15.1P.Seq F M00055804:61 CH15CON 2122 651088 RTA22200014F.c.15.2.P.Seq F M00056945-211 CH16COP 2123 651088 RTA22200014F.c.15.2.P.Seq F M00056945-211 CH16COP 2124 772810 RTA22200014F.c.15.2.P.Seq F M00056945-211 CH16COP 2125 408499 RTA22200014F.c.15.2.P.Seq F M00056945-211 CH16COP 2125 408499 RTA22200007F.d.06.1P.Seq F M00056945-211 CH16COP 2126 556325 RTA22200007F.d.04.1P.Seq F M00056977-88 CH15CON 2129 454776 RTA2220001F.d.07.1P.Seq F M00056977-88 CH15CON 2129 454776 RTA2220001F.d.07.1P.Seq F M00056932:710 CH17COHLV 2130 377679 RTA2220001F.d.07.1P.Seq F M00056932:711 CH16COP 2133 7728131 RTA2220001F.d.07.1P.Seq F M00056932:711 CH17COHLV 2131 728131 RTA2220001F.d.07.1P.Seq F M00056932:711 CH17COHLV 2132 475203 RTA22200007F.d.04.1P.Seq F M00056332:711 CH17COHLV 2133 FX42200007F.d.04.1P.Seq F M00056332:711 CH17COHLV 2133 FX4220001F.d.05.1P.Seq F M00056979-88 CH16COP 2133 3727314 RTA2220001F.d.05.1P.Seq F M00056937-80 CH16COP 2133 3727314 RTA2220001F.d.05.1P.Seq F M00056937-80 CH16COP 2133 772314 RTA2220001F.d.05.1P.Seq F M00056332:711 CH17COHLV 2135 561332 RTA2220001F.d.05.1P.Seq F M0005693-78 CH16COP 2134 45476 RTA2220001F.d.05.1P.Seq F M00056937-80 CH16COP 2134 45420 RTA2220001F.d.05.1P.Seq F M0005693-78 CH16COP 2134 45468 RTA2220001F.d.05.1P.Seq F M0005693-78 CH16COP 2134 45468 RTA2220001F.d.05.1P.Seq F M0005690-75 CH16COP 2144 440488 RTA2220001F.d.05.1P.Seq F M0005690-75 CH16COP 2144 440488 RTA222000	\$					<u>.</u>
2113   55886   RTA22200004F.j0.3.1.P.Seq   F   M00055783:311   CH15CON   CH15CON   2115   454869   RTA22200004F.e.j0.3.1.P.Seq   F   M00055771:44   CH15CON   CH15CON   2116   559674   RTA22200023F.j0.5.1.P.Seq   F   M00055771:42   CH15CON   CH15CON   2117   2235   RTA2220023F.j0.5.1.P.Seq   F   M00055771:42   CH17COHLV   CH17C	Secretary was a consumption of the consumption of t				( Terrorrows & Transactor Commission #58 (Transactor American Amer	2
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2117   2235   RTA2220023F.n.2.1.P.Seq   F   M0002419:31   CH03MAH   2118   7545   RTA2220022F.k.6.6.1.P.Seq   F   M0004054:75   CH01COH   2119   729173   RTA22200016F.n.22.1.P.Seq   F   M00057272:11   CH16COP   2120   650448   RTA22200004F.m.09.1.P.Seq   F   M0005555:0:16   CH15CON   2121   172013   RTA22200014F.c.15.1.P.Seq   F   M0005555:0:16   CH15CON   2122   651088   RTA22200014F.c.15.1.P.Seq   F   M00056945:211   CH16COP   2123   651088   RTA22200014F.c.15.2.P.Seq   F   M00056945:211   CH16COP   2124   726810   RTA22200014F.c.15.2.P.Seq   F   M00056945:211   CH16COP   2125   406499   RTA22200014F.c.15.1.P.Seq   F   M00056947:12   CH15CON   2126   556325   RTA22200024F.i.03.1.P.Seq   F   M00055927:12   CH15CON   2126   556325   RTA22200024F.i.3.1.P.Seq   F   M00055927:12   CH15CON   2127   644836   RTA22200004F.i.3.1.P.Seq   F   M00055927:14   CH17COHLV   2127   644836   RTA22200004F.i.3.1.P.Seq   F   M00055927:48   CH15CON   2128   649062   RTA22200007F.d.0.1.P.Seq   F   M00055927:48   CH16COP   2130   377579   RTA22200025F.f.17.1.P.Seq   F   M000565927:20   CH16COP   2130   377579   RTA22200014F.o.08.1.P.Seq   F   M00056593:212   CH16COP   2131   728131   RTA22200014F.o.08.1.P.Seq   F   M00056593:212   CH16COP   2133   727314   RTA22200014F.o.08.1.P.Seq   F   M00057024:75   CH16COP   2134   552025   RTA22200014F.o.08.1.P.Seq   F   M00057305:62.56   CH15CON   2136   735279   RTA22200014F.o.22.1.P.Seq   F   M00057306:62.56   CH15CON   2136   735279   RTA22200014F.o.08.2.P.Seq   F   M00057306:62.56   CH15CON   2136   735279   RTA22200014F.o.08.2.P.Seq   F   M00057307:812   CH16COP   2137   167   RTA22200014F.o.08.2.P.Seq   F   M00057306:14   CH16COP   2141   724616   RTA22200014F.o.08.2.P.Seq   F   M00057008:31   CH16COP   2141   744616   RTA22200014F.o.08.2.P.Seq   F   M00057008:31   CH16C	Exercise Annual Company	***************************************			tario de la compania	ili manana ana manana ana ana ana ana ana a
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2119   729173	A					
2120   650448   RTA22200004F.m.09.1.P.Seq   F   M000555818:612   CH15CON	Annual commence and the second	5				
2121   172013   RTA2220002F j.16.1.P.Seq   F   M00055520:16   CH15CON   2122   651088   RTA2220014F.c.15.1.P.Seq   F   M00056945:211   CH16COP   2124   726810   RTA22200015F.o.06.1.P.Seq   F   M00056945:211   CH16COP   2125   406499   RTA2220004F.o.13.1.P.Seq   F   M00055827:12   CH15CON   2126   556325   RTA2220002F.0.3.1.P.Seq   F   M00055827:12   CH15CON   2127   644836   RTA2220002F.0.3.1.P.Seq   F   M00055927:48   CH15CON   2128   649062   RTA22200004F.1.3.1.P.Seq   F   M00055797:48   CH15CON   2129   454776   RTA2220001F.d.07.1.P.Seq   F   M00056162:59   CH15CON   2130   377579   RTA22200025F.17.1.P.Seq   F   M0005533:2711   CH17COHLV   2131   728131   RTA2220014F.o.88.1.P.Seq   F   M0005533:2711   CH17COHLV   2132   475203   RTA2220001F.a.05.1.P.Seq   F   M00057024:75   CH16COP   2133   727314   RTA22200014F.o.88.1.P.Seq   F   M00057024:75   CH16COP   2134   552025   RTA22200019F.k.13.1.P.Seq   F   M00056595:78   CH16COP   2134   552025   RTA22200019F.k.13.1.P.Seq   F   M00056595:78   CH16COP   2135   561382   RTA22200014F.o.82.P.Seq   F   M0005679:310   CH17COHLV   2136   732579   RTA22200014F.o.22.1.P.Seq   F   M0005679:525   CH16COP   2137   167   RTA22200014F.o.22.1.P.Seq   F   M0005679:512   CH16COP   2138   185585   RTA22200014F.o.22.1.P.Seq   F   M0005679:512   CH16COP   2139   728131   RTA22200014F.o.22.1.P.Seq   F   M0005679:512   CH16COP   2140   475203   RTA22200014F.o.22.1.P.Seq   F   M00057011:43   CH16COP   2141   724616   RTA22200015F.d.3.1.P.Seq   F   M00057011:43   CH16COP   2142   726594   RTA22200015F.d.3.1.P.Seq   F   M0005502:33   CH16COP   2143   64522   RTA22200016F.d.3.1.P.Seq   F   M00057011:43   CH16COP   2144   400362   RTA22200015F.d.3.1.P.Seq   F   M0005708:39   CH16COP   2145   646583   RTA22200016F.d.3.1.P.Seq   F   M0005708:39   CH16COP   2146   475203   RTA22200005F.d.3.1.P.Seq   F   M0005708:39   CH16COP   2146   475203   RTA22200009F.m.15.1.P.Seq   F   M0005708:39   CH16COP   2146   475203   RTA22200009F.m.15.1.P.Seq   F   M0005500:49   CH16COP   2146   475203   RTA22	Secretary and a construction of the construction of		AND CONTRACTOR OF THE PROPERTY			&100 00 per commence a para commence de commence de para commence de para commence de commence de para commence
2122   651088   RTA22200014F.c.15.1.P.Seq   F   M00056945:211   CH16COP   2123   651088   RTA22200015F.o.06.1.P.Seq   F   M00056945:211   CH16COP   2124   726810   RTA22200015F.o.06.1.P.Seq   F   M0005736.61   CH16COP   2125   406499   RTA22200025F.0.03.1.P.Seq   F   M00055827:12   CH15CON   2126   556325   RTA22200023F.I.03.1.P.Seq   F   M00055087:110   CH17COHLV   2127   644836   RTA22200004F.i.13.1.P.Seq   F   M00055087:110   CH17COHLV   2128   649062   RTA22200007F.d.04.1.P.Seq   F   M00056578.8   CH15CON   2129   454776   RTA22200014F.d.07.1.P.Seq   F   M00056551:62   CH16COP   2130   377679   RTA22200025F.f.17.1.P.Seq   F   M00056551:62   CH16COP   2131   728131   RTA22200014F.o.08.1.P.Seq   F   M00057024:75   CH16COP   2132   475203   RTA22200014F.o.08.1.P.Seq   F   M00057024:75   CH16COP   2133   727314   RTA22200011F.p.15.1.P.Seq   F   M00056597:62   CH16COP   2134   552025   RTA22200014F.a.13.1.P.Seq   F   M00056565:76   CH16COP   2135   561382   RTA22200014F.a.2.1.P.Seq   F   M00056766:25   CH16COP   2136   732679   RTA22200017F.a.22.1.P.Seq   F   M000570300:62   CH16COP   2136   732679   RTA22200017F.a.22.1.P.Seq   F   M000570300:62   CH16COP   2138   185585   RTA22200017F.a.22.1.P.Seq   F   M00057030:62   CH16COP   2138   185585   RTA22200014F.o.02.1.P.Seq   F   M00057030:62   CH16COP   2139   728131   RTA22200014F.o.02.1.P.Seq   F   M00057030:62   CH16COP   2140   475203   RTA22200014F.o.02.1.P.Seq   F   M00057030:63   CH16COP   2140   475203   RTA22200014F.o.02.1.P.Seq   F   M0005701:43   CH16COP   2141   724616   RTA22200014F.o.02.1.P.Seq   F   M0005703:72   CH16COP   2141   724616   RTA22200014F.o.02.1.P.Seq   F   M0005703:73   CH16COP   2142   726594   RTA22200004F.d.13.1.P.Seq   F   M0005703:73   CH16COP   2144   400362   RTA22200004F.d.13.1.P.Seq   F   M0005703:73   CH16COP   2144   400362   RTA22200004F.d.15.1.P.Seq   F   M0005703:75   CH16COP   2144   406362   RTA22200004F.d.15.1.P.Seq   F   M0005503:71   CH15CON   2144   64683   RTA22200004F.d.9.1.P.Seq   F   M0005503:71   CH15CON   214					-	Acres (all all all all all all all all all al
2123   651088   RTA22200014F.c.15.2.P.Seq   F   M00056945:211   CH16COP   2124   726810   RTA22200014F.o.06.1.P.Seq   F   M00057136:61   CH16COP   2125   406499   RTA22200004F.n.23.1.P.Seq   F   M00055087:110   CH17COHLV   2126   556325   RTA22200004F.i.13.1.P.Seq   F   M00055087:110   CH17COHLV   2127   644836   RTA22200004F.i.13.1.P.Seq   F   M00055797:48   CH15CON   2128   649062   RTA22200007F.d.04.1.P.Seq   F   M00056195:79   CH15CON   2129   454776   RTA22200011F.d.07.1.P.Seq   F   M00056551:62   CH15COP   2130   377579   RTA22200025F.f.17.1.P.Seq   F   M00055032:711   CH17COHLV   2131   728131   RTA22200014F.o.08.1.P.Seq   F   M00057024:75   CH16COP   2132   475203   RTA22200016F.a.05.1.P.Seq   F   M00057024:75   CH16COP   2133   727314   RTA22200016F.a.05.1.P.Seq   F   M00057154:24   CH16COP   2134   552025   RTA222200016F.i.3.1.P.Seq   F   M0005659:78   CH16COP   2135   651382   RTA22200004F.d.11.1.P.Seq   F   M0005769:55   CH15CON   2136   732579   RTA22200017F.a.22.1.P.Seq   F   M00057300:62   CH16COP   2137   167   RTA22200012F.o.22.1.P.Seq   F   M00057024:75   CH16COP   2138   185585   RTA22200014F.m.07.2.P.Seq   F   M00056037:28   CH06COP   2140   475203   RTA22200014F.o.08.2.P.Seq   F   M00057024:75   CH16COP   2141   724616   RTA22200014F.o.08.2.P.Seq   F   M00057024:75   CH16COP   2141   724616   RTA22200015F.d.13.1.P.Seq   F   M0005706:14   CH16COP   2141   724616   RTA22200015F.d.13.1.P.Seq   F   M0005706:14   CH16COP   2141   724616   RTA22200015F.d.13.1.P.Seq   F   M0005706:15   CH15CON   2144   400362   RTA22200015F.d.13.1.P.Seq   F   M0005706:15   CH15CON   2145   646583   RTA22200008F.a.16.1.P.Seq   F   M0005708:50   CH16COP   2146   475203   RTA22200008F.a.16.1.P.Seq   F   M0005708:50   CH16COP   2147   550001   RTA22200009F.m.15.1.P.Seq   F   M0005708:50   CH16COP   2148   646583   RTA22200009F.m.15.1.P.Seq   F   M0005708:50   CH16COP   2146   475203   RTA22200009F.m.15.1.P.Seq   F   M00056295:71   CH15CON   2149   646583   RTA22200009F.m.15.1.P.Seq   F   M0005630:15   CH16COP   2150	Samuel Annual Communication of the Communication of	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~				de la companya de la
2124   726810   RTA22200015F.o.06.1.P.Seq   F   M00057136.61   CH16COP   2125   406499   RTA22200004F.n.23.1.P.Seq   F   M00055827.12   CH15CON   2126   556325   RTA22200004F.n.31.P.Seq   F   M00055087.110   CH17COHLV   2127   644836   RTA22200004F.n.31.P.Seq   F   M00055797.48   CH15CON   2128   649062   RTA2220001F.d.04.1.P.Seq   F   M00056597.48   CH15CON   2129   454776   RTA2220001F.d.07.1.P.Seq   F   M00056327.10   CH16COP   2130   377579   RTA22200015F.d.07.1.P.Seq   F   M00056332.711   CH17COHLV   2131   728131   RTA22200014F.o.08.1.P.Seq   F   M00057024.75   CH16COP   2132   475203   RTA22200016F.a.05.1.P.Seq   F   M00056532.71   CH17COHLV   2133   727314   RTA22200011F.p.15.1.P.Seq   F   M00056597.8   CH16COP   2134   555025   RTA22200019F.k.13.1.P.Seq   F   M00056523.110   CH17COHLV   2135   561382   RTA22200014F.o.22.1.P.Seq   F   M00057307.5   CH16COP   2136   732579   RTA22200017F.a.22.1.P.Seq   F   M00057307.5   CH16COP   2137   167   RTA22200012F.o.22.1.P.Seq   F   M00057307.5   CH16COP   2138   185585   RTA222200014F.o.08.2.P.Seq   F   M0005701.5   CH16COP   2140   475203   RTA22200014F.m.08.2.P.Seq   F   M0005701.5   CH16COP   2141   724616   RTA22200014F.m.07.2.P.Seq   F   M0005701.5   CH16COP   2142   726594   RTA22200014F.m.07.2.P.Seq   F   M0005701.5   CH16COP   2143   646583   RTA22200014F.m.07.2.P.Seq   F   M0005701.4   CH16COP   2144   400362   RTA22200014F.m.07.1.P.Seq   F   M0005701.4   CH16COP   2145   646583   RTA22200014F.m.07.1.P.Seq   F   M0005701.4   CH16COP   2146   475203   RTA22200014F.m.07.1.P.Seq   F   M0005701.4   CH16COP   2147   550001   RTA22200014F.m.07.1.P.Seq   F   M0005701.4   CH16COP   2148   640703   RTA22200014F.m.07.1.P.Seq   F   M0005701.4   CH16COP   2149   646583   RTA22200014F.m.07.1.P.Seq   F   M0005501.4   CH16COP   2149   646583   RTA22200014F.m.07.1.P.Seq   F   M0005701.4   CH16COP   2150   449468   RTA22200014F.m.07.1.P.Seq   F   M0005708.5   CH16COP   2151   449468   RTA22200004F.h.07.1.P.Seq   F   M0005501.3   CH16COP   2153   449468   RTA22200004F.	Superior and the second					
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2136         732579         RTA22200017F.a.22.1.P.Seq         F         M00057300:62         CH16COP           2137         167         RTA22200012F.o.22.1.P.Seq         F         M00056779:512         CH16COP           2138         185585         RTA22200241F.I.21.1.P.Seq         F         M00026937:28         CH04MAL           2139         728131         RTA22200014F.o.08.2.P.Seq         F         M00057024:75         CH16COP           2140         475203         RTA22200014F.m.07.2.P.Seq         F         M00057011:43         CH16COP           2141         724616         RTA22200011F.I.16.1.P.Seq         F         M00056622:33         CH16COP           2142         726594         RTA22200015F.d.13.1.P.Seq         F         M00057066:14         CH16COP           2143         645222         RTA22200008F.a.16.1.P.Seq         F         M00056300:15         CH15CON           2144         400362         RTA22200009F.m.15.1.P.Seq         F         M00055222:85         CH17COHLV           2145         646583         RTA22200009F.m.15.1.P.Seq         F         M0005701:43         CH16COP           2147         550001         RTA22200002F.i.9.1.P.Seq         F         M00055454:17         CH15CON           2149         6	<u> </u>					<u> </u>
2137         167         RTA22200012F.o.22.1.P.Seq         F         M00056779:512         CH16COP           2138         185585         RTA22200241F.I.21.1.P.Seq         F         M00026937:28         CH04MAL           2139         728131         RTA22200014F.o.08.2.P.Seq         F         M00057024:75         CH16COP           2140         475203         RTA22200014F.m.07.2.P.Seq         F         M00057011:43         CH16COP           2141         724616         RTA22200011F.I.16.1.P.Seq         F         M00056622:33         CH16COP           2142         726594         RTA222000015F.d.13.1.P.Seq         F         M0005706:14         CH16COP           2143         645222         RTA22200008F.a.16.1.P.Seq         F         M00055030:15         CH15CON           2144         400362         RTA222000024F.j.22.1.P.Seq         F         M0005503:10         CH16COP           2145         646583         RTA22200009F.m.15.1.P.Seq         F         M0005701:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055226:71         CH15CON           2150         44	Supremitation of Merchanical American	ADMITTANCE AND ADMITT			PARTIE AND THE SECOND S	
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2139         728131         RTA22200014F.o.08.2.P.Seq         F         M00057024:75         CH16COP           2140         475203         RTA22200014F.m.07.2.P.Seq         F         M00057011:43         CH16COP           2141         724616         RTA22200011F.l.16.1.P.Seq         F         M00056622:33         CH16COP           2142         726594         RTA22200015F.d.13.1.P.Seq         F         M00057066:14         CH16COP           2143         645222         RTA22200008F.a.16.1.P.Seq         F         M00056300:15         CH15CON           2144         400362         RTA22200024F.j.22.1.P.Seq         F         M00055222:85         CH17COHLV           2145         646583         RTA22200009F.m.15.1.P.Seq         F         M00057011:43         CH16COP           2146         475203         RTA22200014F.m.07.1.P.Seq         F         M00057011:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055203:71         CH15CON           2150         449468         RTA222000014F.l.23.1.P.Seq         F         M0005708:59         CH16COP           2151         <	3	management and the second			M00056779:512	harren anno anno anno anno anno anno anno an
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2141         724616         RTA22200011F.I.16.1.P.Seq         F         M00056622:33         CH16COP           2142         726594         RTA22200015F.d.13.1.P.Seq         F         M00057066:14         CH16COP           2143         645222         RTA22200008F.a.16.1.P.Seq         F         M00056300:15         CH15CON           2144         400362         RTA22200024F.j.22.1.P.Seq         F         M00055222:85         CH17COHLV           2145         646583         RTA22200009F.m.15.1.P.Seq         F         M00055222:85         CH16COP           2146         475203         RTA22200014F.m.07.1.P.Seq         F         M00057011:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M0005708:59         CH16COP           2151         449468         RTA22200014F.l.23.2.P.Seq         F         M0005708:59         CH16COP           2153 <td< td=""><td>Same and the same /td><td></td><td></td><td></td><td>with the contract of the contr</td><td></td></td<>	Same and the same				with the contract of the contr	
2142         726594         RTA22200015F.d.13.1.P.Seq         F         M00057066:14         CH16COP           2143         645222         RTA22200008F.a.16.1.P.Seq         F         M00056300:15         CH15CON           2144         400362         RTA22200024F.j.22.1.P.Seq         F         M00055222:85         CH17COHLV           2145         646583         RTA22200009F.m.15.1.P.Seq         F         M00057011:43         CH16COP           2146         475203         RTA22200002F.e.03.1.P.Seq         F         M00057011:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M00057008:59         CH16COP           2151         449468         RTA22200001F.l.16.1.P.Seq         F         M00057008:59         CH16COP           2153         449468         RTA22200001F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2154	Samuel Sa		<del></del>			}
2143         645222         RTA22200008F.a.16.1.P.Seq         F         M00056300:15         CH15CON           2144         400362         RTA22200024F.j.22.1.P.Seq         F         M00055222:85         CH17COHLV           2145         646583         RTA22200009F.m.15.1.P.Seq         F         M00042850:310         CH16COP           2146         475203         RTA22200014F.m.07.1.P.Seq         F         M00057011:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M00057008:59         CH16COP           2151         449468         RTA22200014F.l.23.2.P.Seq         F         M00042891:34         CH15CON           2152         551628         RTA222000014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA222000021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2156	Samuel	724616			M00056622:33	\$*************************************
2144         400362         RTA22200024F.j.22.1.P.Seq         F         M00055222:85         CH17COHLV           2145         646583         RTA2220009F.m.15.1.P.Seq         F         M00042850:310         CH16COP           2146         475203         RTA22200014F.m.07.1.P.Seq         F         M00057011:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M0005708:59         CH16COP           2151         449468         RTA22200016F.h.22.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA22200014F.l.23.2.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M0005708:59         CH16COP           2154         417259         RTA22200024F.h.07.1.P.Seq         F         M00054804:812         CH17COHLV           2156	A				M00057066:14	CH16COP
2145         646583         RTA22200009F.m.15.1.P.Seq         F         M00042850:310         CH16COP           2146         475203         RTA22200014F.m.07.1.P.Seq         F         M00057011:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M0005708:59         CH16COP           2151         449468         RTA22200014F.l.23.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA22200001F.l.16.1.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M0005708:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157	2143	645222	RTA22200008F.a.16.1.P.Seq	F		
2146         475203         RTA22200014F.m.07.1.P.Seq         F         M00057011:43         CH16COP           2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M00057008:59         CH16COP           2151         449468         RTA22200001F.l.16.1.P.Seq         F         M00057008:59         CH16COP           2152         551628         RTA222000014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2153         449468         RTA222000014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA222000021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA222000024F.h.07.1.P.Seq         F         M00055201:13         CH16COP           2156         524363         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158	2144	400362	RTA22200024F.j.22.1.P.Seq	F	M00055222:85	CH17COHLV
2147         550001         RTA22200002F.e.03.1.P.Seq         F         M00055454:17         CH15CON           2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M00057008:59         CH16COP           2151         449468         RTA22200014F.l.23.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA222000014F.l.23.2.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200024F.h.07.1.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159	2145	646583	RTA22200009F.m.15.1.P.Seq	F	M00042850:310	CH16COP
2148         640703         RTA22200002F.i.19.1.P.Seq         F         M00055512:47         CH15CON           2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M00057008:59         CH16COP           2151         449468         RTA22200016F.h.22.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA22200001F.l.16.1.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00057008:59         CH16COP           2155         448029         RTA22200021F.i.19.3.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00056190:72         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	2146	475203	RTA22200014F.m.07.1.P.Seq	F	M00057011:43	CH16COP
2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M00057008:59         CH16COP           2151         449468         RTA22200016F.h.22.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA22200001F.l.16.1.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200021F.i.19.3.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056190:72         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	2147	550001	RTA22200002F.e.03.1.P.Seq	F	M00055454:17	CH15CON
2149         646583         RTA22200007F.k.10.2.P.Seq         F         M00056236:71         CH15CON           2150         449468         RTA22200014F.l.23.1.P.Seq         F         M00057008:59         CH16COP           2151         449468         RTA22200016F.h.22.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA22200001F.l.16.1.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200021F.i.19.3.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056190:72         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	2148	640703	RTA22200002F.i.19.1.P.Seq	F	M00055512:47	CH15CON
2151         449468         RTA22200016F.h.22.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA22200001F.l.16.1.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200009F.n.10.1.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	2149	646583	RTA22200007F.k.10.2.P.Seq	F	M00056236:71	CH15CON
2151         449468         RTA22200016F.h.22.1.P.Seq         F         M00057223:36         CH16COP           2152         551628         RTA22200001F.l.16.1.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.l.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200009F.n.10.1.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	2150	449468	RTA22200014F.I.23.1.P.Seq		M00057008:59	CH16COP
2152         551628         RTA22200001F.I.16.1.P.Seq         F         M00042891:34         CH15CON           2153         449468         RTA22200014F.I.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200009F.n.10.1.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	2151	449468		F	M00057223:36	
2153         449468         RTA22200014F.I.23.2.P.Seq         F         M00057008:59         CH16COP           2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200009F.n.10.1.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	2152	551628	RTA22200001F.I.16.1.P.Seq	F	M00042891:34	
2154         417259         RTA22200021F.i.19.3.P.Seq         F         M00054804:812         CH17COHLV           2155         448029         RTA22200009F.n.10.1.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	Special and the second	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			M00057008:59	·
2155         448029         RTA22200009F.n.10.1.P.Seq         F         M00042856:67         CH16COP           2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	3 mary mary mary mary mary mary mary mary	garant garant ann an ann an ann an an an an an an an			M00054804:812	~**************************************
2156         524363         RTA22200024F.h.07.1.P.Seq         F         M00055201:13         CH17COHLV           2157         446531         RTA22200023F.n.24.1.P.Seq         F         M00055116:22         CH17COHLV           2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	3		49000000000000000000000000000000000000	taria a construir de la constr	***************************************	
2157       446531       RTA22200023F.n.24.1.P.Seq       F       M00055116:22       CH17COHLV         2158       561359       RTA22200008F.a.11.1.P.Seq       F       M00056295:67       CH15CON         2159       711297       RTA22200007F.f.10.1.P.Seq       F       M00056190:72       CH15CON	\$		CONTRACTOR DE LA CONTRA	i versumman et a como estra que esta esta como esta en como el manerio de la como el manerio del la co	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	CONTRACTOR
2158         561359         RTA22200008F.a.11.1.P.Seq         F         M00056295:67         CH15CON           2159         711297         RTA22200007F.f.10.1.P.Seq         F         M00056190:72         CH15CON	Summerum	processor a reconstruction of the contraction of			~~~~	***************************************
2159 711297 RTA22200007F.f.10.1.P.Seq F M00056190:72 CH15CON	Same			and the second s		and the comment of th
and the contract of the contra	Same and the same	harman and a second	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		~~~	Secretaria de la constante de
		acomentense and a second and a	CONTRACTOR OF THE PROPERTY OF		·	CONTRACTOR

Table 1

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2195	551434	RTA22200003F.m.02.1.P.Seq	F	M00055701:410	CH15CON
2196	727447	RTA22200012F.m.18.1.P.Seq	F	M00056759:73	CH16COP
2197	378786	RTA22200025F.o.11.2.P.Seq	F	M00055396:77	CH17COHLV
2198	649152	RTA22200006F.p.11.2.P.Seq	F	M00056131:312	
2199	18853	RTA22200023F.p.10.1.P.Seq	F	M00055134:53	CH17COHLV
2200	643481	RTA22200005F.f.05.1.P.Seq	F	M00055884:510	CH15CON
2201	644417	RTA22200004F.p.08.1.P.Seq	F	M00055838:712	CH15CON
2202	726788	RTA22200004F.p.06.1.F.Seq	F	M00056597:32	CH16COP
2202	206	RTA22200011F.I.17.1.F.Seq	F	M00030397.32 M00027594:33	\$~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
2203	395930	HENNING CONTRACTOR CON	F	M00056548:311	CH04MAL
2204	185589	RTA22200011F.c.20.1.P.Seq RTA22200244F.a.15.1.P.Seq	F		CH16COP
Successive commence of the commence of	and the second contract of the	учествення по применяющими в применяющими в применяющими в приментории в применяющими в применяющими в применя	F	M00027168:810	CH04MAL
2206	1441	RTA22200225F.i.15.1.P.Seq	F F	M00005485:25	CH02COH
2207	14522	RTA22200241F.h.19.1.P.Seq		M00026909:73	CH04MAL
2208	203605	RTA22200013F.n.02.1.P.Seq	F	M00056896:65	CH16COP
2209	551527	RTA22200019F.g.15.1.P.Seq	F.	M00054496:29	CH17COHLV
2210	4509	RTA22200225F.m.07.1.P.Seq	F	M00005514:16	CH02COH
2211	447737	RTA22200001F.g.12.1.P.Seq	F	M00042704:69	CH15CON
2212	447388	RTA22200004F.k.08.1.P.Seq	<u> </u>	M00055804:61	CH15CON
2213	451932	RTA22200002F.c.24.1.P.Seq	<u> </u>	M00055447:84	CH15CON
2214	559043	RTA22200025F.c.20.1.P.Seq	F	M00055294:44	CH17COHLV

Table 1

	ble 1				
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2215	380634	RTA22200021F.n.06.2.P.Seq	F	M00054842:311	CH17COHLV
2216	495715	RTA22200018F.p.18.1.P.Seq	F	M00043379:37	CH17COHLV
2217	451932	RTA22200002F.d.01.1.P.Seq	F	M00055447:84	CH15CON
2218	447939	RTA22200002F.d.12.1.P.Seq	F	M00055450:79	CH15CON
2219	1181	RTA22200250F.c.03.1.P.Seq	F	M00027781:54	CH04MAL
2220	376600	RTA22200017F.a.06.1.P.Seq	F	M00057288:58	CH16COP
2221	234761	RTA22200001F.j.10.1.P.Seq	F	M00042739:41	CH15CON
2222	644417	RTA22200014F.d.20.1.P.Seq	F	M00056952:16	CH16COP
2223	639048	RTA22200004F.c.05.1.P.Seq	F	M00055749:811	CH15CON
2224	11452	RTA22200226F.d.16.1.P.Seq	F	M00005628:310	CH02COH
2225	452076	RTA22200001F.c.17.1.P.Seq	F	M00042551:49	CH15CON
2226	644523	RTA22200005F.c.07.1.P.Seq	F	M00055864:82	CH15CON
2227	554678	RTA22200021F.h.11.3.P.Seq	F	M00054776:61	CH17COHLV
2228	3550	RTA222000211 .H.11.3.F.Seq	F	M00034770:01	CH03MAH
2229	<u>,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,</u>	RTA22200230F.k.10.1.F.Seq	F	M00056167:28	CH15CON
Samuel Company of the	450311		F	M00055976:61	
2230	647280	RTA22200005F.o.13.1.P.Seq	F		CH15CON
2231	548858	RTA22200026F.f.18.1.P.Seq		M00055480:810	
2232	4204	RTA22200225F.p.22.1.P.Seq	<u> </u>	M00005568:29	CH02COH
2233	540690	RTA22200011F.f.19.1.P.Seq	F _	M00056574:17	CH16COP
2234	404774	RTA22200009F.k.06.2.P.Seq	F	M00042833:71	CH16COP
2235	557823	RTA22200023F.h.01.1.P.Seq	F	M00055057:63	CH17COHLV
2236	1458	RTA22200242F.g.04.1.P.Seq	F	M00027014:74	CH04MAL
2237	485431	RTA22200020F.e.01.1.P.Seq	F	M00054595:89	CH17COHLV
2238	2245	RTA22200229F.I.11.1.P.Seq	F	M00007032:69	CH02COH
2239	3242	RTA22200226F.p.09.1.P.Seq	F	M00005826:710	CH02COH
2240	648747	RTA22200007F.m.12.1.P.Seq	F	M00056254:82	CH15CON
2241	3805	RTA22200222F.c.01.1.P.Seq	F	M00001470:71	CH01COH
2242	475203	RTA22200012F.c.17.1.P.Seg	F	M00056688:65	CH16COP
2243	12018	RTA22200240F.h.11.1.P.Seq	F	M00023399:38	CH04MAL
2244	475203	RTA22200009F.i.18.2.P.Seq	F	M00042818:512	CH16COP
2245	3805	RTA22200222F.b.24.1.P.Seq	F	M00001470:71	CH01COH
2246	496132	RTA22200018F.e.23.1.P.Seq	F	M00043304:41	CH17COHLV
2247	650600	RTA22200005F.m.01.1.P.Seq	F	M00055941:212	CH15CON
2248	650749	RTA22200004F.g.10.1.P.Seq	F	M00055785:63	CH15CON
2249	223148	RTA22200023F.n.05.1.P.Seq	F	M00055106:47	CH17COHLV
2250	449	RTA22200018F.m.12.1.P.Seq	F	M00043355:811	CH17COHLV
2251	735620	RTA22200013F.g.07.1.P.Seq	F	M00056843:89	CH16COP
2252	650600	RTA22200005F.I.24.1.P.Seq	F	M00055941:212	CH15CON
2253	218	RTA22200021F.m.03.2.P.Seq	F	M00054832:59	CH17COHLV
2254	4161	RTA22200222F.d.07.1.P.Seq	F	M00001561:71	CH01COH
2255	373202	RTA22200025F.f.01.1.P.Seq	F F	M00055322:51	CH17COHLV
2256	724339	RTA22200010F.I.14.1.P.Seq	F	M00056490:52	CH16COP
2257	113291	RTA22200007F.p.17.1.P.Seq	F	M00056290:65	CH15CON
2258	736753	RTA22200016F.f.22.1.P.Seq	F	M00057208:48	CH16COP
2259	650600	RTA22200003F.m.18.1.P.Seq	F	M00055706:11	CH15CON
2260	451569	RTA222000031.III.10.11.F.36q	F	M00055700:11	CH16COP
2261	1297	RTA22200013F.n.06.1.P.Seq	F	M00030075:12 M00021655:82	CH03MAH
2262	3680	RTA22200233F.h.00.1.F.Seq	F	M00021033.02 M00026906:310	CH04MAL
	· · · · · · · · · · · · · · · · · · ·	RTA22200005F.g.11.1.P.Seq	F		
2263	63602	angga kanang a paka ang ang kanang kanan	F	M00055888:25	CH15CON
2264	2757	RTA22200238F.I.07.1.P.Seq	5	M00022973:77	CH03MAH
2265	373128	RTA22200020F.a.11.1.P.Seq	E	M00054569:87	CH17COHLV
2266	641479	RTA22200014F.I.16.2.P.Seq	F	M00057007:212	CH16COP
2267	450380	RTA22200015F.b.12.1.P.Seq	<u> </u>	M00057049:85	CH16COP
2268	133512	RTA22200024F.i.12.1.P.Seq	F	M00055209:18	CH17COHLV

Table 1

	ble 1			<del></del>	· <del>- · ·</del>
SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	AND THE RESIDENCE OF THE PROPERTY OF THE PROPE	LIBRARY
2269	447211	RTA22200006F.b.19.2.P.Seq	F	M00056007:66	CH15CON
2270	645222	RTA22200013F.a.24.1.P.Seq	F	M00056802:81	CH16COP
2271	645222	RTA22200013F.b.01.1.P.Seq	F	M00056802:81	CH16COP
2272	17372	RTA22200002F.i.22.1.P.Seq	F	M00055513:46	CH15CON
2273	451619	RTA22200025F.m.23.2.P.Seq	F	M00055388:88	CH17COHLV
2274	2510	RTA22200238F.m.21.1.P.Seq	F	M00022995:52	CH03MAH
2275	643974	RTA22200007F.n.15.1.P.Seq	F	M00056268:24	CH15CON
2276	500630	RTA22200013F.j.19.1.P.Seq	F	M00056872:16	CH16COP
2277	3101	RTA22200225F.n.05.1.P.Seq	F	M00005530:17	CH02COH
2278	446938	RTA22200004F.c.11.1.P.Seq	F	M00055751:41	CH15CON
2279	554469	RTA22200013F.f.23.1.P.Seq	F	M00056841:79	CH16COP
2280	554469	RTA22200003F.n.24.1.P.Seq	F	M00055721:66	CH15CON
2281	2894	RTA22200016F.b.05.1.P.Seq	F	M00057163:21	CH16COP
2282	650600	RTA22200008F.e.04.1.P.Seq	F	M00056338:210	CH15CON
2283	3101	RTA22200226F.j.12.1.P.Seq	F	M00005710:41	CH02COH
2284	554469	RTA22200003F.o.01.1.P.Seq	F	M00055721:66	CH15CON
2285	9910	RTA22200236F.n.06.1.P.Seq	F	M00022667:35	CH03MAH
2286	400608	RTA22200008F.g.03.1.P.Seq	F	M00056460:711	CH15CON
2287	555051	RTA22200021F.o.07.2.P.Seg	F	M00054854:58	CH17COHLV
2288	185400	RTA22200002F.j.21.1.P.Seq	F	M00055522:57	CH15CON
2289	3059	RTA22200233F.g.05.1.P.Seq	F	M00008075:112	CH03MAH
2290	647185	RTA22200004F.b.20.1.P.Seq	F	M00055747:49	CH15CON
2291	1669	RTA22200237F.p.19.1.P.Seq	F	M00022853:311	CH03MAH
2292	7158	RTA22200232F.I.21.1.P.Seq	F	M00022129:512	CH03MAH
2293	496132	RTA22200005F.d.06.1.P.Seq	F	M00055871:76	CH15CON
2294	378623	RTA22200024F.k.20.1.P.Seq	F	M00055227:52	CH17COHLV
2295	1257	RTA22200227F.i.05.1.P.Seq	F	M00006679:34	CH02COH
2296	648499	RTA22200012F.d.21.1.P.Seq	F	M00056698:512	CH16COP
2297	185627	RTA22200242F.k.21.1.P.Seq	F	M00027042:711	CH04MAL
2298	640005	RTA22200008F.c.15.1.P.Seq	F	M00056312:33	CH15CON
2299	553462	RTA22200021F.e.14.3.P.Seg	F	M00054750:412	CH17COHLV
2300	649852	RTA22200002F.g.01.1.P.Seq	F	M00055470:71	CH15CON
2301	422375	RTA22200018F.e.04.1.P.Seq	F	M00043296:210	CH17COHLV
2302	10910	RTA22200227F.n.22.1.P.Seq	F	M00006745:42	CH02COH
2303	2737	RTA22200023F.c.21.1.P.Seq	F	M00055034:71	CH17COHLV
2304	3438	RTA22200225F.o.18.1.P.Seq	F	M00005548:12	CH02COH
2305	3438	RTA22200228F.m.18.2.P.Seq	F	M00006885:711	CH02COH
2306	3763	RTA22200225F.o.12.1.P.Sea	F	M00005546:72	CH02COH
2307	648966	RTA22200007F.a.18.1.P.Seq	F	M00056141:29	CH15CON
2308	724339	RTA22200013F.m.10.1.P.Seq	F	M00056892:11	CH16COP
2309	451569	RTA22200018F.I.17.1.P.Seq	F	M00043350:86	CH17COHLV
2310	554109	RTA22200007F.d.17.1.P.Seq	F	M00056171:812	CH15CON
2311	380339	RTA22200006F.h.10.2.P.Seq	F	M00056055:17	CH15CON
2312	729903	RTA22200014F.g.06.1.P.Seq	F	M00056966:111	CH16COP
2313	45	RTA22200025F.I.10.1.P.Seq	F	M00055379:38	CH17COHLV
2314	454653	RTA22200019F.d.21.1.P.Seq	F	M00043503:55	CH17COHLV
2315	11536	RTA222000131.d.21.111.Seq	F	M00043003:36	CH03MAH
2316	373134	RTA22200001F.h.02.1.P.Seq	F	M00042711:34	CH15CON
2317	185691	RTA22200041F.g.04.1.P.Seq	F	M00032711:34 M00026900:32	CH04MAL
2318	234761	RTA22200011F.o.06.1.P.Seq	F	M00056646:75	CH16COP
2319	724339	RTA222000117 .0.00:117 .0eq	F	M00056710:31	CH16COP
2320	732740	RTA22200012F.i.19.11F.Seq	F	M00057231:19	CH16COP
2321	35895	RTA22200010F.d.10.1.P.Seq	F	M0005725119	CH16COP
2322	133512	RTA222000177.d.10.1.F.Seq	F	M00037323.370	CH17COHLV
2022	100012	1111 EEEE000 101 .0.10.11.1 .004		11100072011.01	OTT TOOLIEV

Table 1

1 8	ble 1				
SEQ ID	MARKON AMERICANA OROMANIAN SERVICE	SEQ NAME	ORIENTATION	CONTROLOGICA CONTR	LIBRARY
2323	2974	RTA22200227F.b.03.1.P.Seq	F	M00006590:310	CH02COH
2324	500	RTA22200231F.I.13.1.P.Seq	F	M00007983:43	CH03MAH
2325	376919	RTA22200020F.m.09.1.P.Seq	F	M00054678:57	CH17COHLV
2326	8403	RTA22200233F.g.08.1.P.Seq	F	M00008076:62	CH03MAH
2327	3643	RTA22200227F.e.11.1.P.Seq	F	M00006638:72	CH02COH
2328	447211	RTA22200001F.j.13.1.P.Seq	F	M00042741:410	CH15CON
2329	447211	RTA22200001F.b.15.1.P.Seq	F	M00042538:18	CH15CON
2330	14929	RTA22200025F.i.20.1.P.Seq	F	M00055363:712	CH17COHLV
2331	648934	RTA22200006F.k.14.2.P.Seq	F	M00056081:29	CH15CON
2332	731785	RTA22200014F.j.11.2.P.Seq	F	M00056993:43	CH16COP
2333	639908	RTA22200005F.g.08.1.P.Seq	F	M00055887:54	CH15CON
2334	344577	RTA22200014F.m.20.1.P.Seq	F	M00057015:312	CH16COP
2335	2906	RTA22200015F.m.13.1.P.Seq	F	M00057127:611	CH16COP
2336	446938	RTA22200002F.c.18.1.P.Seq	F	M00055445:76	CH15CON
2337	2493	RTA22200234F.h.08.1.P.Seq	F	M00022251:19	CH03MAH
2338	38	RTA22200006F.o.08.2.P.Seq	F	M00056112:82	CH15CON
2339	13818	RTA22200238F.e.13.1.P.Seq	F	M00022898:64	CH03MAH
2340	8371	RTA22200229F.o.12.1.P.Seq	F	M00007065:611	CH02COH
2341	402494	RTA22200012F.o.16.1.P.Seq	F	M00056776:16	CH16COP
2342	731785	RTA22200014F.j.11.1.P.Seq	F	M00056993:43	CH16COP
2343	4621	RTA22200235F.f.15.1.P.Seq	F	M00022468:510	CH03MAH
2344	9750	RTA22200229F.g.23.1.P.Seq	F	M00006979:210	CH02COH
2345	133512	RTA22200024F.p.23.1.P.Seq	F	M00055263:79	CH17COHLV
2346	162626	RTA22200250F.f.03.1.P.Seq	F	M00027823:77	CH04MAL
2347	730059	RTA22200011F.a.20.1.P.Seq	F	M00056530:41	CH16COP
2348	2069	RTA22200224F.k.12.1.P.Seq	F	M00005373:86	CH02COH
2349	5868	RTA22200242F.i.17.1.P.Seq	F F	M00027030:38	CH04MAL
2350	2683	RTA22200226F.o.08.1.P.Seq	F	M00005813:510	CH02COH
2351	380409	RTA22200012F.h.07.1.P.Seq	F	M00056719:612	CH16COP
2352	639991	RTA22200002F.k.20.1.P.Seq	F	M00055527:54	CH15CON
2353	535	RTA22200233F.a.19.1.P.Seq	F	M00008015:210	CH03MAH
2354	14929	RTA22200025F.f.21.1.P.Seq	F	M00055335:83	CH17COHLV
2355	134702	RTA22200249F.n.23.1.P.Seq	F	M00027733:45	CH04MAL
2356	642477	RTA22200004F.j.16.1.P.Seq	F	M00055802:612	CH15CON
2357	14929	RTA22200047 .j. 10.117 .Seq	F	M00055002:012	CH16COP
2358	134702	RTA222000147.i.13.2.F.Seq	F	M00036950:011	CH04MAL
2359	185649	RTA22200250F.g.08.1.P.Seq	F	M00020931:70	CH04MAL
2360	10702	RTA222002301 .g.06.1.F .Seq	F	M00027853:41 M00026860:51	CH04MAL
	643955	RTA22200001F.k.21.1.P.Seq	F	M00020800:31	CH15CON
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2362	643955		F	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	CH03MAH
2363	4455 195567	RTA22200233F.o.08.1.P.Seq	F	M00021670:75	
2364	185567	RTA22200242F.c.16.1.P.Seq	F	M00026985:25	CH04MAL
2365	9115	RTA22200226F.e.04.1.P.Seq	F	M00005632:82	CH16COR
2366	14929	RTA22200015F.i.12.1.P.Seq	F	M00057100:59	CH16COP
2367	14929	RTA22200014F.i.15.1.P.Seq		M00056986:611	CH16COP
2368	4181	RTA22200244F.k.18.1.P.Seq	F	M00027203:88	CH04MAL
2369	5206	RTA22200238F.n.19.1.P.Seq	<u> </u>	M00023002:710	CH03MAH
2370	825	RTA22200237F.c.05.1.P.Seq	<u> </u>	M00022702:24	CH03MAH
2371	825	RTA22200238F.o.24.1.P.Seq	<u> </u>	M00023020:79	CH03MAH
2372	825	RTA22200231F.d.09.1.P.Seq	F	M00007943:32	CH03MAH
2373	2748	RTA22200225F.d.18.1.P.Seq	<u> </u>	M00005449:510	CH02COH
2374	2748	RTA22200238F.i.02.1.P.Seq	<u> </u>	M00022934:23	CH03MAH
2375	2748	RTA22200225F.d.08.1.P.Seq	<u> </u>	M00005445:12	CH02COH
2376	133512	RTA22200014F.a.15.2.P.Seq	F	M00056923:59	CH16COP

Table 1

SEQ ID	CLUSTER	SEQ NAME	ORIENTATION	CLONE ID	LIBRARY
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2378	642477	RTA22200008F.e.02.1.P.Seq	F	M00056338:26	CH15CON
2379	642477	RTA22200007F.n.16.1.P.Seq	F	M00056268:41	CH15CON
2380	2493	RTA22200236F.f.17.1.P.Seq	F	M00022624:32	CH03MAH
2381	5796	RTA22200235F.j.03.2.P.Seq	F	M00022508:32	CH03MAH
2382	3782	RTA22200236F.k.02.1.P.Seq	F	M00022655:410	CH03MAH
2383	884	RTA22200016F.b.01.1.P.Seq	F	M00057162:82	CH16COP
2384	5860	RTA22200238F.p.04.1.P.Seq	F	M00023023:63	CH03MAH
2385	5275	RTA22200226F.h.09.1.P.Seq	F	M00005675:72	CH02COH
2386	3932	RTA22200233F.I.12.1.P.Seq	F	M00021626:34	CH03MAH
2387	884	RTA22200017F.a.17.1.P.Seq	F	M00057297:63	CH16COP
2388	4455	RTA22200235F.j.23.2.P.Seq	F	M00022517:53	CH03MAH
2389	5860	RTA22200236F.c.15.1.P.Seq	F	M00022598:45	CH03MAH
2390	5860	RTA22200232F.m.23.1.P.Seq	F	M00022143:310	CH03MAH
2391	372791	RTA22200003F.o.02.1.P.Seq	F	M00055721:55	CH15CON
2392	5206	RTA22200238F.o.14.1.P.Seq	F	M00023015:410	CH03MAH
2393	372791	RTA22200005F.o.07.1.P.Seq	F	M00055973:44	CH15CON
2394	2846	RTA22200228F.I.12.2.P.Seq	F	M00006873:512	CH02COH
2395	5275	RTA22200225F.g.17.1.P.Seq	F	M00005469:17	CH02COH
2396	2846	RTA22200228F.i.21.2.P.Seq	F	M00006852:37	CH02COH

	ACCESSION	DESCRIPTION1	P VALU
63	3236249	(AC004684) hypothetical protein [Arabidopsis thaliana]	9.9
		H-2 CLASS II HISTOCOMPATIBILITY ANTIGEN, GAMMA	
		CHAIN (HLA-DR ANTIGENS ASSOCIATED INVARIANT	-
		CHAIN) >gi 92086 pir  S04362 class II histocompatibility antigen-	
64	123111	associated gamma chain, long splice form - rat	9.7
65	94377	hypothetical protein 221 - turnip yellow mosaic virus	9.6
		(AF039047) contains similarity to the BPTI/kunitz family of	1
66	2736449	inhibitors [Caenorhabditis elegans]	9.5
67	543894	BETA-LACTAMASE PRECURSOR	8.2
		(Z70038) cDNA EST EMBL:D32579 comes from this gene; cDNA	
		EST EMBL:D35254 comes from this gene; cDNA EST yk224b3.5	
68	3881525	comes from this gene; cDNA EST yk357f10.5 comes from this gene	7.6
69	3128358	(AF010496) ribose transport system permease protein RbsC	7.5
		(Z68297) Weak similarity to Mouse DNA-binding protein BMI-1	·
		(SW:BMI1_MOUSE); cDNA EST EMBL:C07407 comes from this	
70	3875771	gene; cDNA EST EMBL:C07408 comes from this gene	7.3
*******	21293	(Y00759) 20 kDa protein (AA 1-212) [Spinacia oleracea]	7.3
72	2131007	(Z95890) pknE [Mycobacterium tuberculosis]	7.2
	245923	(S83583) multiple-epitope polypeptide 1, MEP-1 construct]	7
MARION CONTRACTOR STATES	tananii kultaaniin 1990aan olehaan ole	PROCOLLAGEN ALPHA 2(IV) CHAIN PRECURSOR	
		>gi 84486 pir  \$16366 collagen alpha 2(IV) chain precursor - pig	
74	115347	roundworm >gi 159649 (M67507) putative [Ascaris suum]	6.9
	2072674	(Z95120) rhlE [Mycobacterium tuberculosis]	5.8
	807646	น้อง การแรงการทางการและการทางการการทางการการการการการการการการการการการการการก	5.8
***************************************		URIDINE 5'-MONOPHOSPHATE SYNTHASE OROTIDINE 5'-	
77	131706	PHOSPHATE DECARBOXYLASE	5.6
	631593	glucose transport protein homolog - sheep	5
The second secon	854064	(X83413) U87 [Human herpesvirus 6]	4.8
80	484695	vascular cell adhesion molecule 1 - human	4.2
81	1236146	(U49864) fus1 protein [Chlamydomonas reinhardtii]	3.8
		acetyl-CoA carboxylase (EC 6.4.1.2) - Cyclotella cryptica >gi 409450	ź
82	1083846	(L20784) acetyl-CoA carboxylase [Cyclotella cryptica]	3.5
83	2245054	(Z97342) protein kinase homolog	3.4
		UBIQUITIN-CONJUGATING ENZYME E2-21.2 KD (UBIQUITIN	<del>}</del>
		PROTEIN LIGASE) (UBIQUITIN CARRIER PROTEIN)	
84	1717863	>gi 1077331 pir  S51438 probable membrane protein YLR306w -	3.4
85	1363331	transcription factor IIIC alpha chain - rat alpha-subunit [Rattus	3.4
86	2736327	(AF038615) No definition line found [Caenorhabditis elegans]	3.1
87	4587895	(AF072509) glutamate receptor interacting protein 2 [Rattus	3.1
mandermonare many said	1480746	(U62529) matrix metalloproteinase 3 [Equus caballus]	3
	225858	thyroid/steroid receptor related gene [Homo sapiens]	3
***************************************	3861293		2.4
~~~~			2
processore and the second	3256583		1.7
······································	1684985	(U20633) NADH dehydrogenase subunit [Neuwiedia veratrifolia]	1.5
	452517	(D26361) KIAA0042 [Homo sapiens]	1.4
	4218558		1.1
**************************************	4539280	(AL049498) putative transcription factor	0.28
~~~~	4539280	(AL049498) putative transcription factor	0.26
		(Z78201) Similarity to E.coli 2-oxoglutarate dehydrogenase	
Anamounge		(SW:ODO1_ECOLI); cDNA EST EMBL:D32590 comes from this	
Management		gene; cDNA EST EMBL:D32841 comes from this gene; cDNA EST	
98	3881842	EMBL:D34051 comes from this gene; cDNA EST EMBL:D35268	4e-019
		,,	.~ (1)

100 33270 102 31269 103 39307 106 27087 107 73355 108 38517 112 21207 120 33345 121 24079  122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 302500 139 38818	79 76 41 4 03 29 26 56 01 80 38 48 94	(AB014524) KIAA0624 protein [Homo sapiens] (AF062483) SDP3 [Homo sapiens] (AF099149) TRIAD1 type I [Homo sapiens] (AC003952) hypothetical protein [Arabidopsis thaliana] (U23450) similar to RNA-binding protein [Caenorhabditis elegans] (AF100421) p80 [Rattus norvegicus] GumG protein - Xanthomonas campestris (AL021306) predicted using FGENEH [Homo sapiens] (X87612) mono ATP-ribosaltransferase [Mus musculus] (Z69662) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail MINOR CAPSID PROTEIN L2 >gi 1020224 type 36] HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this gene; cDNA EST EMBL:C13538 comes from this gene; cDNA EST	
103 39307 106 27087 107 73355 108 38517 112 21207 120 33345 121 24079 123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818	76 41 4 03 29 26 56 56 01 80 38 48 94	(AF099149) TRIAD1 type I [Homo sapiens]  (AC003952) hypothetical protein [Arabidopsis thaliana]  (U23450) similar to RNA-binding protein [Caenorhabditis elegans]  (AF100421) p80 [Rattus norvegicus]  GumG protein - Xanthomonas campestris  (AL021306) predicted using FGENEH [Homo sapiens]  (X87612) mono ATP-ribosaltransferase [Mus musculus]  (Z69662) predicted using Genefinder; similar to collagen; cDNA  EST EMBL:D75049 comes from this gene; cDNA EST  EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223  comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail  MINOR CAPSID PROTEIN L2 >gi 1020224 type 36]  HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN  (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w),  Hypothetical protein, len: 430 aa  HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE  INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical  (Z79759) Similarity to Yeast endosomal P24A protein  (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	9e-068 3.6 1e-014 7e-055 6 7.7 5.9 5.9 3.5 3.4 3.4 0.61
106 27087 107 73355 108 38517 112 21207 120 33345 121 24079 122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818 140 194549 141 37460	41 4 03 29 26 56 01 80 38 48 94	(AC003952) hypothetical protein [Arabidopsis thaliana] (U23450) similar to RNA-binding protein [Caenorhabditis elegans] (AF100421) p80 [Rattus norvegicus] GumG protein - Xanthomonas campestris (AL021306) predicted using FGENEH [Homo sapiens] (X87612) mono ATP-ribosaltransferase [Mus musculus] (Z69662) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail MINOR CAPSID PROTEIN L2 >gi 1020224 type 36] HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	3.6 1e-014 7e-055 6 7.7 5.9 3.5 3.4 3.4 3.4 9.8
107 73355 108 38517 112 21207 120 33345 121 24079  122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 302500 139 38818	4 03 29 26 56 56 01 80 38 48 94	(U23450) similar to RNA-binding protein [Caenorhabditis elegans]     (AF100421) p80 [Rattus norvegicus]     GumG protein - Xanthomonas campestris     (AL021306) predicted using FGENEH [Homo sapiens]     (X87612) mono ATP-ribosaltransferase [Mus musculus]     (Z69662) predicted using Genefinder; similar to collagen; cDNA     EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223     comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail     MINOR CAPSID PROTEIN L2 > gi 1020224 type 36     HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii > gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN     (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w),     Hypothetical protein, len: 430 aa     HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE     INTERGENIC REGION > gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	1e-014 7e-055 6 7.7 5.9 5.9 3.5 3.4 3.4 0.61
108 38517 112 21207 120 33345 121 24079  122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818	03 29 26 56 01 80 38 48 94	[(AF100421) p80 [Rattus norvegicus]  [GumG protein - Xanthomonas campestris [(AL021306) predicted using FGENEH [Homo sapiens] [(X87612) mono ATP-ribosaltransferase [Mus musculus] [(Z69662) predicted using Genefinder; similar to collagen; cDNA [EST EMBL:D75049 comes from this gene; cDNA EST [EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST EMBL:D72768 comes from this [Comes from this gene; cDNA EST CEMSB40F comes from this [Comes from this gene; cDNA EST CEMSB40F comes from this [Comes from this gene; cDNA EST CEMSB40F comes from this [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Comes from this gene; cDNA EST CEMSB40F comes from this] [Com	7e-055 6 7.7 5.9 3.5 3.4 3.4 0.61
112 21207 120 33345 121 24079 122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818: 140 194549 141 37460	29 26 56 01 80 38 48 94	GumG protein - Xanthomonas campestris  (AL021306) predicted using FGENEH [Homo sapiens]  (X87612) mono ATP-ribosaltransferase [Mus musculus]  (Z69662) predicted using Genefinder; similar to collagen; cDNA  EST EMBL:D75049 comes from this gene; cDNA EST  EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223  comes from this gene; cDNA EST EMBL:D72768 comes from this  serine/threonine protein kinase - quail  MINOR CAPSID PROTEIN L2 >gi 1020224 type 36]  HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii  >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus  NEUROGENIC LOCUS NOTCH 3 PROTEIN  (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w),  Hypothetical protein, len: 430 aa  HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE  INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical  (Z79759) Similarity to Yeast endosomal P24A protein  (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	5.9 3.5 3.4 3.4 0.61
120 33345 121 24079 122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818 140 19454 141 37460	26 56 56 01 80 38 48 94	(AL021306) predicted using FGENEH [Homo sapiens] (X87612) mono ATP-ribosaltransferase [Mus musculus] (Z69662) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail MINOR CAPSID PROTEIN L2 >gi 1020224 type 36] HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	7.7 5.9 3.5 3.4 3.4 0.61
121 24079  122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 302506 139 38818	56 01 80 38 48 94 71	(X87612) mono ATP-ribosaltransferase [Mus musculus] (Z69662) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail MINOR CAPSID PROTEIN L2 >gi 1020224 type 36] HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	7.7 5.9 3.5 3.4 3.4 0.61
122 38777 123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818 140 194549 141 37460	01 80 38 48 94 71	(X87612) mono ATP-ribosaltransferase [Mus musculus] (Z69662) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail MINOR CAPSID PROTEIN L2 >gi 1020224 type 36] HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	3.5 3.4 3.4 3.4 9.8
123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818 140 19454 141 37460	80 38 48 94 71	(Z69662) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail  MINOR CAPSID PROTEIN L2 >gi 1020224 type 36]  HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN  (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa  HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE  INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein  (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	3.5 3.4 3.4 3.4 9.8
123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818 140 19454 141 37460	80 38 48 94 71	EST EMBL:D75049 comes from this gene; cDNA EST EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail  MINOR CAPSID PROTEIN L2 >gi 1020224 type 36]  HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN  (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa  HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein  (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	3.5 3.4 3.4 3.4 0.61 9.8
123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818 140 19454 141 37460	80 38 48 94 71	EMBL:D72067 comes from this gene; cDNA EST EMBL:D72223 comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail  MINOR CAPSID PROTEIN L2 >gi 1020224 type 36]  HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN  (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa  HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE  INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein  (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	3.5 3.4 3.4 3.4 0.61 9.8
123 21177 124 17227 125 24944 126 24942 127 44939 138 30250 139 38818 140 19454 141 37460	80 38 48 94 71	comes from this gene; cDNA EST EMBL:D72768 comes from this serine/threonine protein kinase - quail  MINOR CAPSID PROTEIN L2 >gi 1020224 type 36]  HYPOTHETICAL PROTEIN MJ0208 Methanococcus jannaschii >gi 1498983 (U67476) 4Fe-4S iron-sulfur protein [Methanococcus NEUROGENIC LOCUS NOTCH 3 PROTEIN  (AL034559) predicted using hexExon; MAL3P7.11 (PFC0910w), Hypothetical protein, len: 430 aa  HYPOTHETICAL 28.8 KD PROTEIN IN MOAE-RHLE  INTERGENIC REGION >gi 1787008 (AE000181) orf, hypothetical (Z79759) Similarity to Yeast endosomal P24A protein  (SW:EM70_YEAST); cDNA EST CEMSB40F comes from this	3.5 3.4 3.4 3.4 0.61 9.8
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140 194549 141 37460		gene; cDNA EST EMBL:C13538 comes from this gene: cDNA EST	
140 194549 141 37460		gene: CDNA EST EMBL:C13538 comes from this gene: cDNA FST	4
141 37460	, ,	(IISCOCS) Circilanta MADON	9.6
141 37460		(U56965) Similar to NAD(P) transhydrogenase, mitochondrial; coded	d
141 37460		for by C. elegans cDNA yk27c1.5; coded for by C. elegans cDNA	Management
141 37460		yk35b9.5; coded for by C. elegans cDNA yk35b9.3; coded for by C.	-
~~~~~	13	elegans cDNA yk161c9.3; coded for by C. elegans	5.6
		(AC005311) putative GTP-binding protein [Arabidopsis thaliana]	5.6
142 208884	.3	(AF003386) F59E12.9 gene product [Caenorhabditis elegans]	3.2
		GLUCAN ENDO-1,3-BETA-GLUCOSIDASE PRECURSOR ((1-	
143 170655	1	>3)-BETA-GLUCAN ENDOHYDROLASE) aestivum]	1.9
144 629778		chitinase (EC 3.2.1.14) - barley vulgare]	1.9
145 279127	······	(Z95327) Cleavage Stimulation Factor sapiens]	0.85
146 736767	recorrection and a second	helper component protease [Turnip mosaic virus]	0.5
W. W		HYPOTHETICAL 35.4 KD PROTEIN CY20G9.19C	·\$
147 173120		>gi 1449291 emb CAB00954  (Z77162) hypothetical protein	0.22
148 387706	3	(Z37092) F44F4.10 [Caenorhabditis elegans]	0.075
- Commission		(Z73428) similar to Zinc finger, C3HC4 type (RING finger); cDNA	1
	]	EST EMBL:D67323 comes from this gene [Caenorhabditis elegans]	***************************************
APRICA APPRICA	27	>gi 3881096 emb CAB16481  finger); cDNA EST EMBL:D67323	4
149 387873	9 (	comes from this gene	2e-029
159 445448		(AC006234) putative kinase, 5' partial	9.1
160 304408	6 (	[AF055904) unknown [Myxococcus xanthus]	5.4
***************************************	) [	U56965) Similar to NAD(P) transhydrogenase, mitochondrial; coded	J. <b>T</b>
***		For by C. elegans cDNA yk27c1.5; coded for by C. elegans cDNA	
	1	rk35b9.5; coded for by C. elegans cDNA yk35b9.3; coded for by C.	
161 194549	3 =	elegans cDNA yk161c9.3; coded for by C. elegans	E 1
162 730883		COLUMN SET OF COURT OF DY C. Clegans	5.4
102 / 50003	······································	SVNAPTIC VESICIE DDOTEINIO (CVO)	
163 235121	S	SYNAPTIC VESICLE PROTEIN 2 (SV2) norvegicus] D88386) ggg pol polygrotain (propygorania) [Figinal 1]	5.4
コロッシスペコンコ	<u> </u>	SYNAPTIC VESICLE PROTEIN 2 (SV2) norvegicus] D88386) gag-pol polyprotein (precursor protein) [Friend murine eukemia virus]	5.4 4.2

		or (BlastX vs. Non-Redundant Proteins)	
SEQ ID	ACCESSION	DESCRIPTION1	P VALUE
	***************************************	(AF118222) contains similarity to ubiquitin carboxyl-terminal	
		hydrolase family 2 (Pfam:PF00443, score=48.3, E=3.5e-13, N=2)	ALE SANGERS
165	4115922	and (Pfam:PF00442, Score=40.0 E=5.2e-08, N=1) [Arabidopsis	2.4
		VERY HYPOTHETICAL 52.7 KD PROTEIN C8A4.05C IN	
166	1351639	CHROMOSOME I >gi 2130446 pir  S62521 hypothetical protein	2.4
		small inducible cytokine subfamily D (Cys-X3-Cys), member 1	The second
	1	(fractalkine, neurotactin) >gi 1888523 (U84487) CX3C chemokine	
\$	4506857	precursor [Homo sapiens] >gi 1899259 (U91835) CX3C chemokine	2.4
168	4505637	protocadherin 8; PCDH8 sapiens]	1.9
processor and a second a second and a second a second and	3249055	(AF071210) casein kinase II alpha subunit [Spodoptera frugiperda]	1.4
***************************************	854065	(X83413) U88 [Human herpesvirus 6]	1e-005
171	854065	(X83413) U88 [Human herpesvirus 6]	1e-005
191	3150072	(AF046996) preS1 surface protein [woolly monkey hepatitis B Virus	1 6.9
192	2622845	(AE000928) corrinoid/iron-sulfur protein, large subunit	6.9
193	1083554	tyrosine phosphoprotein SLP-76 - mouse	6.9
		LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED	
	* *	PROTEIN PRECURSOR (LRP) Caenorhabditis elegans >gi 156360	***************************************
	50.00	(M96150) LDL receptor-related protein [Caenorhabditis elegans]	********
194	1708868	Genefinder; Identity to C.elegans Low density lipid (LDL) receptor-	5.2
		(AL023702) putative insertion element IS1647 transposase	
195	3169030	[Streptomyces coelicolor]	4
196	728835	!!!! ALU SUBFAMILY SC WARNING ENTRY	4
197		vascular cell adhesion molecule 1 - human	3.9
198		(Y13898) glutathione-S-transferase	3.9
***************************************		(U41554) coded for by C. elegans cDNA yk38a7.3; coded for by C.	13.9
		elegans cDNA yk8c6.3; coded for by C. elegans cDNA yk25d12.5;	
		coded for by C. elegans cDNA yk25d12.3; coded for by C. elegans	*
199	1118071	cDNA yk8c6.5; coded for by C. elegans cDNA yk7f8.5;	2.3
~~~~~	·····	(U24495) 2a protein [Broad bean mottle virus]	1.4
Same and the second	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	E2 protein - human papillomavirus type 18 papillomavirus type 18]	1.4
····	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(L14933) convertase PC5 [Rattus norvegicus]	0.45
***************************************	***************************************	TRANS-ACTING TRANSCRIPTIONAL PROTEIN ICP4	0.73
į		(TRANSCRIPTIONAL ACTIVATOR IE175) (ALPHA-4	
***************************************		PROTEIN) human herpesvirus 1 >gi 59558 emb CAA32286	
Super-dental	o o o o o o o o o o o o o o o o o o o	(X14112) RS1 RS1 [human herpesvirus 1]	
203	124141	>gi 59849 emb CAA29763  1298) [human herpesvirus 1]	0.031
		UNKNOWN >gi 3264861 (U97670) eukaryotic translation initiation	0.031
204	4503511	factor eIF3, p35 subunit [Homo sapiens]	2 - 010
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	UNKNOWN >gi 3264861 (U97670) eukaryotic translation initiation	3e-010
205		factor eIF3, p35 subunit [Homo sapiens]	2- 010
	and the contract of the contra	( A TO SEE C. S. T.	3e-010
	·····	AL023290) putative ATP-dependent RNA helicase	8e-033
221		0.1.	0.7
		A 0007047311	8.7
	1086982 I	1 4 1 · · · · · · · · · · · · · · · · ·	8.7
one comments and the property of	CONTRACTOR AND THE PROPERTY OF	T. ( C.	8.6
	······································	HYPOTHETICAL 433.2 KD PROTEIN IN HXT5-NRK1	8.6
Mineraling		NTERGENIC REGION >gi 626646 pir  S46715 hypothetical protein	* Water
225	731689	THR 000m - yearst (Sanaharamyana ai-i> : 1407000 (Trans and	0.5
<del>สพา</del> จภาคมากการทำ	2351132		8.5
			8.5
	1,32737 (	AT 034550) predicted using hovEven. MALARY 47 (DEC1000)	8.5
228	1493995 H	AL034559) predicted using hexExon; MAL3P7.47 (PFC1080c), Typothetical protein, len: 232 aa	
	F-7-2-7-2	турошенсат ртоещ, теп: 434 aa	6.7

SEQ ID	ACCESSION	bor (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	DATATE
-011X 11D	1 Tree Education	NUCLEOCAPSID PROTEIN coronavirus (strain K378)	P VALU
220	462685	>gi 58850 emb CAA47246	
***************************************	0 4519268	(AB024314) CREA [Aspergillus aculeatus]	5.2
*****	1 1364138		5
	1704138	probable polymerase - soybean dwarf virus polymerase [Soybean	5
	4000	ADENYLOSUCCINATE SYNTHETASE adenylosuccinate synthas	e
222	1709938	(EC 6.3.4.4) - Thiobacillus ferrooxidans >gi 48168 emb CAA40593	\$
232	1709938	(X57324) purA adenylosuccinate synthetase [Thiobacillus	3.8
222	1947132	(AF000262) the second exon has similarity to collagen alpha in a	
***************************************	and the commence of the commence and the commence and the commence of the commence and the	glycine- and proline-rich region	2.9
	2245054	(Z97342) protein kinase homolog	2.9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2736517	(AF039052) contains similarity to helicases	1.7
*************************	1620170	(U42580) a499L [Paramecium bursaria Chlorella virus 1]	1.7
237	4336718	(AF104031) caudal-related homeobox protein	1.3
•••		BETA-GLUCOSIDASE A (GENTIOBIASE) 3.2.1.21) -	
238	114969	Caldocellum saccharolyticum 1-455) [Caldicellulosiruptor	0.76
		HOMEOBOX PROTEIN CDX-1 (CAUDAL-TYPE HOMEOBOX	
239	1170313	PROTEIN 1) >gi 1083361 pir  A49303 homeotic protein Cdx-1 -	0.44
	·	(U44834) polyphosphate glucokinase [Mycobacterium tuberculosis]	
	1172222	>gi 1588398 prf  2208389A phosphate glucokinase [Mycobacterium	8.4
	2959862	(AJ002303) synaptogyrin 1c [Homo sapiens]	8.1
	2983552	(AE000721) cation efflux system (czcD-like) [Aquifex aeolicus]	8
261	104800	nicotinic acetylcholine receptor alpha-5 chain precursor - chicken	4.8
	Managanagag	(Z78013) predicted using Genefinder; cDNA EST EMBL:D72806	***************************************
		comes from this gene; cDNA EST EMBL:D75743 comes from this	
		gene; cDNA EST yk417b6.3 comes from this gene; cDNA EST	
262	3875957	yk417b6.5 comes from this gene; cDNA EST yk276c6.3 com	4.8
263	746475	(U23511) No definition line found [Caenorhabditis elegans]	4.7
264	1899232	(U90333) aquarius gene product [Mus musculus]	2.8
265	3123186	AFLATOXIN BIOSYNTHESIS REGULATORY PROTEIN	2.8
3	\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$\$	NUCLEOLYSIN TIAR (TIA-1 RELATED PROTEIN) >gi 1592563	4.0
266	2500589	(U55861) RNA binding protein TIAR	2.8
267	90253	leukocyte common antigen precursor - mouse musculus]	1.7
268	2827553	(AL021635) predicted protein [Arabidopsis thaliana]	1.6
	3695397	(AF096372) No definition line found [Arabidopsis thaliana]	0.42
	2564330	(AB006629) KIAA0291 [Homo sapiens]	8e-006
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3413860	(AB007918) KIAA0449 protein [Homo sapiens]	3e-009
******		(AF070661) HSPC005 [Homo sapiens]	
	· · · · · · · · · · · · · · · · · · ·	PROBABLE PROCESSING AND TRANSPORT PROTEIN	1e-010
289		(INFECTED CELL PROTEIN 18.5) murine cytomegalovirus (strain	6
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	~~~~~~~~~	(Y13898) glutathione-S-transferase	6
····		(AJ002140) DNA	3.5
resourcement and a second	4567223	The state of the s	3.5
and the same and t	2077849		3.1
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(X 100000)	9.9
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	PROBABLE DEOCESSING AND TRANSPORT PROMERY	7.3
305		PROBABLE PROCESSING AND TRANSPORT PROTEIN	
······································			5.6
····	<b></b>	/3.7.4.0.0.0.\ .1	5.6
30/		(Y13898) glutathione-S-transferase	3.3
***************************************	u	HOMOSERINE KINASE (HK) Pseudomonas aeruginosa	
200	105004	>gi 45425 emb CAA46169  (X65034) homoserine kinase	
·····		Pseudomonas aeruginosa]	3.3
·		(AF015560) RO11 [Neurospora crassa]	0.66
310 4	484695	vascular cell adhesion molecule 1 - human	0.64

SEQ ID	ACCESSION	DESCRIPTION1	P VALU
	***************************************	(Z72502) Similarity with proline-rich proteoglycan (PIR accession	
		number B48013); cDNA EST EMBL:D66054 comes from this gene;	
		cDNA EST EMBL:D69700 comes from this gene; cDNA EST	
311	3874146	yk446b10.3 comes from this gene; cDNA EST yk446b10	0.52
211	3674140	CIRCUMSPOROZOITE PROTEIN PRECURSOR precursor -	0.52
312	117589	Plasmodium knowlesi (strain Nuri) >gi 160198	9e-005
	2781381	(AC004013) Similar to rabbit A-kinase-anchoring protein sapiens]	1e-019
313	2/01301	PAN1 PROTEIN >gi 626783 pir  S48440 poly(A)-specific	10-019
217	720260	ribonuclease (EC 3.1.13.4) - yeast PAN1_YEAST P32521 PAB-	7.0
~~~~	730269	DEPENDENT POLY(A)-SPECIFIC RIBONONUCLEASE but	7.2
	3201610	(AC004669) unknown protein [Arabidopsis thaliana]	5.4
····	3129975	(AL023516) Complement C4 [Gallus gallus]	2.4
322	1850592	(U88295) carnitine palmitoyltransferase II [Rattus norvegicus]	6.9
		(Z70757) Weak similarity to the XFIN protein (Swiss Prot accession	T WE WANTED TO THE TOTAL TOTAL TO THE TOTAL
*****	3881650	number P08045)	6.6
324	2653998	(AF032884) tnsC [Thiobacillus ferrooxidans]	8.2
		(M18083) periplasmic [NiFe]hydrogenase protein small subunit	
326	145111	precursor [Desulfovibrio gigas]	0.37
327	2244839	(Z97337) hypothetical protein [Arabidopsis thaliana]	3.1
328	3935180	(AC004557) F17L21.23 [Arabidopsis thaliana]	4.6
332	3328899	(AE001320) hypothetical protein [Chlamydia trachomatis]	7.4
**********************	4096264	(U26528) Ig heavy chain [Oryctolagus cuniculus]	7.1
		168 KD SURFACE-LAYER PROTEIN PRECURSOR	<u>.</u>
		[CONTAINS: 120 KD SURFACE-EXPOSED PROTEIN	
		MEMBRANE PROTEIN OMPB); 32 KD BETA PEPTIDE]	
334	3023209	precursor - Rickettsia typhi >gi 1871223 (L04661) crystalline surface	5.0
***************************************	4589400	(AB009958) polyprotein [satsuma dwarf virus]	5.7
	7307700	SODIUM-DEPENDENT NORADRENALINE TRANSPORTER	3.1
		(NOREPINEPHRINE TRANSPORTER) (NET) bovine	
226	1709356	>gi 1050439 emb CAA55645  (X79015) norepinephrine transporter	5.4
<del>~~~</del> ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4512671	(AC006931) unknown protein [Arabidopsis thaliana]	1.9
	3757516	(AC005167) putative TMV resistance protein [Arabidopsis thaliana]	3.8
343	3/3/310	HYPOTHETICAL 62.6 KD PROTEIN IN CDS1-RPL2	3.0
244	506400	INTERGENIC REGION >gi 626496 pir  S45886 hypothetical protein	0.00
344	586480		0.26
2.45	100001	88K E-26-specific domain protein Pok - Drosophila	
· sexe consecution and	423981	>gi 217342 dbj BAA01080  (D10228) Ets domain protein	0.054
rando en la companya de la companya	1653153		5.5
348	2147342	E4 protein - human papillomavirus type 14D	0.67
		GENOME POLYPROTEIN [CONTAINS: N-TERMINAL	
		PROTEIN (P1); HELPER COMPONENT PROTEINASE	
and the second		INCLUSION PROTEIN (CI); 6 KD PROTEIN 2 (6K2); GENOME-	
349	3914412	LINKED PROTEIN (VPG); NUCLEA mosaic virus]	0.95
350	2997741	(AF054838) tetraspan TM4SF; Tspan-1 [Homo sapiens]	1e-017
351	1086900	(U41278) contains similarity to G beta repeats	3e-027
		2-OXOGLUTARATE DEHYDROGENASE E1 COMPONENT	
		(ALPHA-KETOGLUTARATE DEHYDROGENASE)	
372	129036	dehydrogenase [Azotobacter vinelandii]	9.8
		cellulose synthase - Agrobacterium tumefaciens >gi 710493 (L38609)	
373	2120777		9.7
arramana ar	628527		9.6
2,7		HOMEOBOX PROTEIN NK-1 (S59/2) fly (Drosophila	
375	123395		7.8
	3722000	(AF035323) survival motor neuron protein	********* *****************************
2/0	J144000	(ATOSSSES) SULVIVAL INDIOL HUMON DIOLEM	7.8

SEQ ID	ACCESSION	DESCRIPTION1	P VALUE
377	3881030	(AL021493) Y51A2B.2 [Caenorhabditis elegans]	7.4
378	4505777	PHD finger protein 1 >gi 2660720 1) [Homo sapiens]	6
379	2190501	(X53706) immunoglobulin alpha-2 heavy chain [Pan troglodytes]	5.8
		(AE001171) conserved hypothetical integral membrane protein	<b>/</b>
380	2688643	[Borrelia burgdorferi]	5.7
**************************************		(U41264) coded for by C. elegans cDNA cm13g1; Similar to	***************************************
381	1086832	bumetanide-sensitive Na-K-Cl cotransporter.	5.4
ipo	1931639	(U95973) lysophospholipase isolog [Arabidopsis thaliana]	4.6
	3757516	(AC005167) putative TMV resistance protein [Arabidopsis thaliana]	4.5
···		interferon consensus sequence binding protein 1 >gi 2275153	
384	4504567	(M91196) DNA-binding protein [Homo sapiens]	4.4
		SUR1 PROTEIN >gi 542362 pir  S41798 SUR1 protein - yeast	<u> </u>
385	464822	(Saccharomyces cerevisiae) cerevisiae] >gi 976268 dbj BAA05628	4.4
	***************************************	EXOSTOSIN-L (MULTIPLE EXOSTOSIS-LIKE PROTEIN)	
386	3023738	>gi 1524413 (U67191) multiple exostosis-like protein [Homo	4.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3878603	(Z83116) M01B2.3 [Caenorhabditis elegans]	3.4
anamana amanana amanana anana an	3745858		3.3
MANAGEMENT OF THE SECONDARIAN	4321758	(AF060669) polyprotein [Hepatitis E virus]	3.3
and the statement of th	2661037	(AF035285) dihydroxyacetone phosphate acyltransferase	3.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2950355	(AJ223300) homebox protein DRx [Drosophila melanogaster]	2.7
	3821973	(AF061140) merozoite surface protein 1 [Plasmodium falciparum]	2.5
and the second s	1552187	(D84375) ORF3 [Oryzias latipes]	.2
373	1332107	S100 CALCIUM-BINDING PROTEIN A13 calcium-binding protein	
304	2493417	A13 [Mus musculus]	1.6
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3550082	(AF071186) WW domain binding protein 11 [Mus musculus]	1.5
		oligodendrocyte-specific proline-rich protein 2 - human	1. <i>J</i> ;
306	1082665	>gi 1408050 dbj BAA05660  (D28114) MOBP [Homo sapiens]	1.5
270	1002005	(Z68296) Similarity to Mouse A-RAF proto-oncogene	1.5
A Company		serine/threonine-protein kinase gene; cDNA EST EMBL:T01018	
307	3874925	comes from this gene; cDNA EST EMBL:D33256 comes from this	1.2
	3014723	PROBABLE SIGNAL RECOGNITION 54 KD PROTEIN (SRP54)	<u> </u>
		>gi 2129283 pir  E64312 signal recognition particle protein -	
308	2500886	Methanococcus jannaschii subunit SRP54 [Methanococcus	1.2
376 2	2300000	(AF016428) contains similarity to Vaccinia virus 37 kd envelope	1.4
300	2291241	protein [Caenorhabditis elegans]	1.1
3972	2271271	(L06798) class D tetracycline/H+ antiporter [Plasmid pRA1]	1.1
400	309958	>gi 575937 dbj BAA03719  (D16172) PP-TETA protein	0.91
<i>พระพรสสสสสสสสสสสสสสสสสสสสสสสสสสสสสสสสสส</i>	3877379	(Z46267) F49E2.2 [Caenorhabditis elegans]	0.39
reconstruction and the second	4539280	(AL049498) putative transcription factor	0.39
<del>~~</del> ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3293235	(U96413) putative transcription factor (U96413) putative opine synthase [Agrobacterium tumefaciens]	0.29
······································	3329636	(AF078786) No definition line found [Caenorhabditis elegans]	0.097
707.	/J#/UJU	(AL023828) cDNA EST yk491f8.5 comes from this gene	
405	3947614	[Caenorhabditis elegans]	8e-019
700	<b>ッノマ / U 1 寸</b>	(AL023828) cDNA EST yk491f8.5 comes from this gene	00-019
406	3947614	[Caenorhabditis elegans]	10.010
400 3	<b>,ノ</b> オ / ひ1十	(AL021481) similar to Phosphoglucomutase and	1e-019
		phosphomannomutase phosphoserine; cDNA EST EMBL:D36168	
***		. · · · · · · · · · · · · · · · · · · ·	
407 3	999030	comes from this gene; cDNA EST EMBL:D70697 comes from this	1 000
~ <del>~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ </del>	3880930	gene; cDNA EST yk373h9.5 comes from this gene; cDNA EST	1e-022
408 3	3184082		1e-028
400	1500550	PUTATIVE RIBONUCLEASE III (RNASE III)	<b></b>
409 2	2500558	>gi 3876420 emb CAB03005  (Z81070) similar to ribonuclease	7e-030

SEQ ID	ACCESSION	DESCRIPTION1	P VALU
	***************************************	(Z48583) similar to ATPases associated with various cellular	7 M 674. MAL . MAL
	No.	activities (AAA); cDNA EST EMBL:Z14623 comes from this gene;	Voltani (america
	Anna Anna Anna Anna Anna Anna Anna Anna	cDNA EST EMBL:D75090 comes from this gene; cDNA EST	N. C.
410	3877493	EMBL:D72255 comes from this gene; cDNA EST yk200e4.5	5e-044
	3298605	(AF057365) UDP N-acetylglucosamine transporter [Canis familiaris]	1e-051
territo di la distribui di tributa e consi		(AF026031) putative mitochondrial outer membrane protein import	
412	4103604	receptor [Homo sapiens]	1e-059
		(AB007902) HH0712 cDNA clone for KIAA0442 has a 574-bp	<del></del>
413	2662165	insertion at position 1474 of the sequence of KIAA0442. [Homo	4e-070
	3882145	(AB018255) KIAA0712 protein [Homo sapiens]	9e-072
	***************************************	CHAPERONE PROTEIN CS3-1 PRECURSOR Escherichia coli	<u> </u>
		>gi 41156 emb CAA34815  (X16944) 27kD protein (AA 1 to 241)	
477	117552	[Escherichia coli]	9.3
	2983622	(AE000726) hypothetical protein [Aquifex aeolicus]	9.3
	2088694	(AF003135) W03F11.1 gene product [Caenorhabditis elegans]	9.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4240237	(AB020681) KIAA0874 protein [Homo sapiens]	9.2
	1864073	(U63002) T-cell receptor beta chain [Callithrix jacchus]	9
.01		DNA helicase II 70K chain homolog - Rhipicephalus appendiculatus	
		>gi 1063592 (L41356) ku autoantigen p70 homologue [Rhipicephalus	
482	2147334	appendiculatus]	8.9
	3323145	(p. 5-5	8.7
***************************************	2183251	(AF002227) putative polyprotein [border disease virus strain C413]	8.6
101	2103231	(Z92830) cDNA EST yk223c7.5 comes from this gene; cDNA EST	0.0
		yk307b2.5 comes from this gene; cDNA EST yk377h2.5 comes from	
		this gene; cDNA EST yk223c7.3 comes from this gene; cDNA EST	
185	3875768	yk307b2.3 comes from this gene [Caenorhabditi	8.6
702	3673700	HYPOTHETICAL PROTEIN MG456 Mycoplasma genitalium	0.0
		(SGC3) >gi 1046175 (U39732) M. genitalium predicted coding	
		region MG456 [Mycoplasma genitalium] >gi 3845050 (U39727)	
186	1351589	conserved hypothetical protein [Mycoplasma genitalium]	8.4
	3023800	GLUCOSE-6-PHOSPHATASE (G6PASE) familiaris]	6.9
	450722	(X71982) ORF j11R [African swine fever virus]	6.8
anno anno anno anno anno anno a	2633756	(Z99111) similar to heavy metal-transporting ATPase [Bacillus	6.6
hananaanaanaan	4049887	(AF063866) ORF MSV024 ALI motif gene family protein	6.5
770	4043007	abdominal-A homeodomain protein - Junonia coenia >gi 797277	0.5
401	1363994		5.3
***************************************	2370493		5.3
anceas conservation accessories of	140631	(Z98944) hypothetical protein HYPOTHETICAL 25.9 KD PROTEIN FP25.9 [Fowlpox virus]	5.3
***************************************	400927		
·····		``````````````````````````````````````	5.1
493	2462935		5
		(AC000132) Similar to A. thaliana receptor-like protein kinase (gb RLK5 ARATH). ESTs gb ATTS0475,gb ATTS4362 come from	
406	2160190	(gb RLR3_ARATH). ESTS gb ATTS04/3,gb ATTS4362 come from this gene. [Arabidopsis thaliana]	_
	2160189	<del>promoteria de la composition della composition </del>	4.0
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1850972		4.9
498	2145678	B1549_C3_230 protein - Mycobacterium leprae	4.2
		CGMP-INHIBITED 3',5'-CYCLIC PHOSPHODIESTERASE A	
400	2192052	(CYCLIC GMP INHIBITED PHOSPHODIESTERASE A) (CGI-	4.0
499	3182952		4.2
<b>5</b> 00	4402520	(AL034358) predicted using hexExon; L4830.1, Hypothetical	
500	4493738		4.2
African		PUTATIVE POLYKETIDE SYNTHASE PKSL 2.3.1) - Bacillus	
		subtilis >gi 40058 emb CAA78479  subtilis] >gi 528996 (U11039)	
	730336	polyketide synthase polyketide synthase of type I [Bacillus subtilis]	4.2
502	188908	(M19720) L-myc protein [Homo sapiens]	4

SEQ ID	ACCESSION	DESCRIPTION1	P VALU
503	4056436	(AC005990) EST gb AA650912 comes from this gene.	3.9
		(Z81035) predicted using Genefinder; Similarity to Sheep	***************************************
504	3874349	vasopressin V1A receptor (SW:P48043)	3.9
505	3874067	(Z93374) similar to 7TM receptor elegans]	3.9
	······	(Z49967) cDNA EST EMBL:T00743 comes from this gene; cDNA	
arawan na		EST EMBL:D69356 comes from this gene; cDNA EST	-
***************************************		EMBL:D65790 comes from this gene; cDNA EST EMBL:D70463	and the same
506	3979818	comes from this gene; cDNA EST EMBL:D66620 comes from this	3.2
	***************************************	ENVELOPE POLYPROTEIN GP160 PRECURSOR 2]	
507	119462	>gi 225570 prf  1306388H gene env [Human immunodeficiency virus	3.2
		COLICIN V PRODUCTION PROTEIN (DEDE PROTEIN) (PUR	
- Annabarra		REGULON 18 KD PROTEIN) colicin V production [Escherichia	-
508	117657	coli] dedE protein [Escherichia coli]	3.1
	***************************************	ATP SYNTHASE A CHAIN (PROTEIN 6) 3.6.1.34) protein 6 - fruit	
		fly (Drosophila yakuba) mitochondrion (SGC4)	
509	114441	>gi 12921 emb CAA25442  (X00924) ATPase subunit 6 [Drosophila	3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1280094	(U55369) No definition line found [Caenorhabditis elegans]	3
200.000.000.000.000.000.000.000.000.000	226131	thyroid hormone receptor alpha 2 [Rattus norvegicus]	3
	<b></b>	(AJ235271) GUANOSINE PENTAPHOSPHATE	1
512	3860855	PHOSPHOHYDROLASE (gppA) [Rickettsia prowazekii]	2.9
312		CELLULOSE SYNTHASE OPERON C PROTEIN xylinus	12.7
513	584834	>gi 1090660 prf  2019362C acsC gene	2.4
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	629777	chitinase (EC 3.2.1.14) - barley vulgare]	2.2
<del>~~~~</del> ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2555183	(AF026504) SPA-1 like protein p1294 [Rattus norvegicus]	2.2
	2333103	forkhead (Drosophila) homolog 1 HEAD DOMAIN PROTEIN	<u> </u>
idia vale		FKHR >gi 631145 pir  S40521 FKHR protein - human >gi 435423	§
		(U02310) fork head domain protein [Homo sapiens]	AA. ( 1000000
516	4503737	>gi 737918 prf  1923399A FKHR gene [Homo sapiens]	1.8
~~~~	3790719	(AF099916) contains similarity to C2H2-type zinc fingers	1.8
	1785942	(U83412) CAG [Drosophila melanogaster]	1.8
- 210	17000772	(Z81079) predicted using Genefinder; similar to collagen; cDNA	1.0
		EST EMBL:M88890 comes from this gene; cDNA EST	-
		EMBL:Z14325 comes from this gene; cDNA EST EMBL:D27520	s -
510	3877036	comes from this gene; cDNA EST EMBL:D72240 comes from this	1.4
317	3677030	hypothetical protein 24 - Agrobacterium tumefaciens plasmid	1.4
520	281654	pTi15955 >gi 39086 emb CAA25186  tumefaciens]	1.3
320	201034	BETA-GLUCOSIDASE (GENTIOBIASE) Ruminococcus albus	1.3
***************************************		>gi 45968 emb CAA33461  (X15415) beta-glucosidase (AA 1 - 947)	aria di dia di
521	114972	[Ruminococcus albus] albus]	1.2
	1707085	(U80451) Similar to collagen [Caenorhabditis elegans]	1.3
JZZ .	1707065	nonstructural protein - hepatitis E virus RNA-directed RNA	1.3
522	102162		1.2
	483163 4455275	polymerase [Hepatitis E virus] (AI 035527) putative protein [Arabidopsis thelional	1.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4455275 4376875	(AL035527) putative protein [Arabidopsis thaliana]	0.99
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4376875	(AE001642) CT465 hypothetical protein	0.45
<i>ขางเทพ</i> พ <b>งและเ</b> คระเทศ (การการการการการการการการการการการการการก	2494911 4520280	HYPOTHETICAL PROTEIN KIAA0124 product is novel. [Homo	0.44
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4539280 1842255	(AL049498) putative transcription factor	0.28
328	1842255	(U74613) hepatocyte nuclear factor-3/fork head homolog 11B [Homo	0.28
500	221077	D(4) DOPAMINE RECEPTOR (D(2C) DOPAMINE RECEPTOR)	0.073
emenumina-anamaning	231977	>gi 203916 (M84009) dopamine receptor D4	0.073
530 9	987050	(X65335) lacZ [Cloning vector pSV-beta-Galactosidase Control]	0.042
	3	S100 CALCIUM-BINDING PROTEIN A13 calcium-binding protein	
varantaria de la constanta de	2493416	A13 (S100A13) [Homo sapiens]	0.031
532	namanamanahamanahan kananah salah salah salah sa serat da se	(U83086) LimA [Dictyostelium discoideum] (AJ224360) surf5c [Homo sapiens]	0.0006
F-0-01-			5e-015

SEQ ID	ACCESSION	.)	P VALL
		(AL023828) cDNA EST yk491f8.5 comes from this gene	
534	3947614	[Caenorhabditis elegans]	1e-015
535	1086860	(U41272) Similar to man(9)-alpha-mannosidase.	3e-028
		(Z66496) cDNA EST EMBL:D71941 comes from this gene; cDNA	1
		EST EMBL:D74691 comes from this gene; cDNA EST	1
		EMBL:D76330 comes from this gene; cDNA EST EMBL:D65192	
536	3875451	comes from this gene; cDNA EST EMBL:D68540 comes from this	2e-030
		(Z48583) similar to ATPases associated with various cellular	<u> </u>
		activities (AAA); cDNA EST EMBL:Z14623 comes from this gene;	
		cDNA EST EMBL:D75090 comes from this gene; cDNA EST	
537	3877493	EMBL:D72255 comes from this gene; cDNA EST yk200e4.5	le-035
538	3169010	(AJ006412) putative GTP-binding protein	2e-042
		(Z48583) similar to ATPases associated with various cellular	1
Y sa alaba		activities (AAA); cDNA EST EMBL:Z14623 comes from this gene;	
		cDNA EST EMBL:D75090 comes from this gene; cDNA EST	
539	3877493	EMBL:D72255 comes from this gene; cDNA EST yk200e4.5	4e-044
		(Z48583) similar to ATPases associated with various cellular	70-044
4		activities (AAA); cDNA EST EMBL:Z14623 comes from this gene;	
COLUMN TO SERVICE SERV		cDNA EST EMBL:D75090 comes from this gene; cDNA EST	***************************************
540	3877493	EMBL:D72255 comes from this gene; cDNA EST yk200e4.5	3e-044
	3882189	(AB018277) KIAA0734 protein [Homo sapiens]	9.9
		(Z48716) similarity to a transmembranous region of ubiquinol-	7.7
ar away away		cytochrome-C reductase (PIR accession number S38960); cDNA	
200		EST EMBL:T00461 comes from this gene; cDNA EST	\$ 2
600	3877937	EMBL:D27071 comes from this gene; cDNA EST EMBL:D27070	9.6
an an arter a commence and a	3643019	(AF064703) glucose transporter 1; CeGT1 [Drosophila	n fr 2.2.0.00
010	3043019	HYPOTHETICAL PROTEIN MJ1394 Methanococcus jannaschii	8.4
611	3219946	,	0
011	3219940	>gi 1592041 (U67579) conserved hypothetical protein HYPOTHETICAL PROTEIN MJ1394 Methanococcus jannaschii	8
612	3219946	>gi 1592041 (U67579) conserved hypothetical protein	o
<del>~~~~</del>	2833328	FIBRILLARIN	8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4505481	<u> </u>	7.9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	220578	nucleoporin 88kD complex protein [Homo sapiens]	7.8
013	220376	(D00570) open reading frame (251 AA) [Mus musculus] NAD(P) TRANSHYDROGENASE SUBUNIT BETA	7.8
- Control of the Cont		transhydrogenase [Escherichia coli] transhydrogenase (B-specific)	1
(1)	266010	(EC 1.6.1.1) b chain NAD(P)+ transhydrogenase (B-specific) (EC	
······································	266810	1.6.1.1) b chain [Escherichia coli] >gi 1787886 (AE000255) pyridine	Karananan x
	807646	(M17294) unknown protein [Human herpesvirus 4]	7.6
618	829186	(X03879) rudimentary protein fragment	7.4
(10	4222246	(AF081825) sodium-dependent high-affinity dicarboxylate	
uma marana manana manana ka	4322346	transporter [Rattus norvegicus]	7.4
620 .	3334785	(AL031107) hypothetical protein SC5A7.04c	7.4
		PHOSPHATIDYLINOSITOL-4-PHOSPHATE 5-KINASE TYPE II	an consequence and
		ALPHA (PIP5KII-ALPHA) KINASE) >gi 1079474 pir  A55967 1-	*
		phosphatidylinositol-4-phosphate 5-kinase (EC 2.7.1.68) - human	
aranan arana a	1346720	>gi 758697 (U14957) 53K isoform of Type II phosphatidylinositol-4-	\$~~~~~
arana ar	4105819		6.4
623	155865	(M93125) 80 kDa protein [Babesia bovis]	6.3
OUT COMPANY		boule protein - fruit fly (Drosophila melanogaster) >gi 1395211	
<del></del>	2133638	(U51858) boule protein	6.2
······································	1788052	(AE000270) putative transport system permease protein	6.2
	3875616	(Z77657) F08H9.9 [Caenorhabditis elegans]	6.2
	2499150	HYPOTHETICAL PROTEIN IN CPS REGION	6.2

		oor (BlastX vs. Non-Redundant Proteins)	
SEQ ID	ACCESSION	DESCRIPTION1	P VALU
		GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2	
		ANTIGEN) (IGE-BINDING PROTEIN) (35 KD LECTIN)	
628	1170758	(CARBOHYDRATE BINDING PROTEIN 35) (CBP 35)	6.1
	***	HEAT SHOCK PROTEIN 42 (42 KD HEAT SHOCK PROTEIN)	
		>gi 1077219 pir  S49767 heat shock protein HSP42 - yeast	
629	2495335	(Saccharomyces cerevisiae)	6
		(U41020) coded for by C. elegans cDNA yk64f5.3; coded for by C.	
630	1086677	elegans cDNA yk64f5.5; Similar to zinc finger. [Caenorhabditis	6
		GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2	***************************************
		ANTIGEN) (IGE-BINDING PROTEIN) (35 KD LECTIN)	- water
***********	1170758	(CARBOHYDRATE BINDING PROTEIN 35) (CBP 35)	6
*****************	440957	Achaete-Scute homolog Mash-1 gene product	6
633	1786037	(U72284) NADH dehydrogenase subunit 2 [Apis mellifera]	6
	delign	HEAT SHOCK PROTEIN 42 (42 KD HEAT SHOCK PROTEIN)	
	A CONTRACTOR OF THE CONTRACTOR	>gi 1077219 pir  S49767 heat shock protein HSP42 - yeast	The same of
634	2495335	(Saccharomyces cerevisiae)	6
		(AF036687) contains similarity to protease inhibitors, WAP-type four	*-
635	2662541	disulfide core domains and thyroglobulin type-1 repeats	5.7
		DAUGHTERLESS PROTEIN fly (Drosophila melanogaster)	1
		>gi 7839 emb CAA68368  melanogaster] >gi 157174 (J03148)	
636	118249	daughterless protein	5.6
637	2447066	(U42580) A570L [Paramecium bursaria Chlorella virus 1]	5.6
638	400927	RIBONUCLEOPROTEIN RB97D ribonucleoprotein [Drosophila	4.9
639	121189	GLUCOSE INHIBITED DIVISION PROTEIN A gidA	4.8
		(AC004238) putative phosphoribosylaminoimidazolecarboxamide	<u> </u>
640	3033398	formyltransferase [Arabidopsis thaliana]	4.6
		POSSIBLE MALTASE PRECURSOR (LARVAL VISCERAL	
		PROTEIN D) >gi 103222 pir  S08597 hypothetical protein D - fruit	NA CONTRACTOR OF THE CONTRACTO
641	126691	fly (Drosophila melanogaster) melanogaster]	4.5
		RETROTRANSPOSABLE ELEMENT SLACS 132 KD PROTEIN	\$6×4×0×00×00×00×00×0×0×0×0×0×0×0×0×0×0×0×
		(ORF2) >gi 84054 pir  S14916 hypothetical protein 2 - Trypanosoma	s 5
642	134087	brucei gambiense transposon SLACS >gi 10535 emb CAA34931	4.3
		(L03172) This CDS feature is included to show the translation of the	·
		corresponding V_region. Presently translation qualifiers on V region	
643	567166	features are illegal.	3.7
644	4355	(Z14126) RhoNUC protein [Saccharomyces cerevisiae]	3.7
645	462679	MYOSIN IB HEAVY CHAIN heavy chain [Dictyostelium	3.6
	######################################	(U41508) similar to C. elegans proteins C26E6.9A and C26E6.9B;	A
646	1098989		2.1
647	3249559		2
648	2435594		2
	995808	(U32240) Ig heavy chain [Mus musculus]	2
650	3281870	(AL031004) putative protein [Arabidopsis thaliana]	1.8
651	728837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	1.6
***************************************	1628461		1.2
052		hypothetical protein - Marek's disease virus gammaherpesvirus	
032	1		
052			
032		tumorigenicity associated mRNA, two complete cds's.], gene products	
CHAPTER AND	Vertrant Westernammen	tumorigenicity associated mRNA, two complete cds's.], gene products [Gallid herpesvirus type 1] >gi 299459 bbs 129316 (S58431) CD4	n 03
CHAPTER AND	483165	tumorigenicity associated mRNA, two complete cds's.], gene products [Gallid herpesvirus type 1] >gi 299459 bbs 129316 (S58431) CD4 precursor homolog to CD4 and IgM heavy chain [Gallid herpesvir	0.93
CHAPTER AND	483165	tumorigenicity associated mRNA, two complete cds's.], gene products [Gallid herpesvirus type 1] >gi 299459 bbs 129316 (S58431) CD4 precursor homolog to CD4 and IgM heavy chain [Gallid herpesvir hypothetical protein - Marek's disease virus gammaherpesvirus	0.93
CHAPTER AND	483165	tumorigenicity associated mRNA, two complete cds's.], gene products [Gallid herpesvirus type 1] >gi 299459 bbs 129316 (S58431) CD4 precursor homolog to CD4 and IgM heavy chain [Gallid herpesvir hypothetical protein - Marek's disease virus gammaherpesvirus tumorigenicity associated mRNA, two complete cds's.], gene products	0.93
653	483165	tumorigenicity associated mRNA, two complete cds's.], gene products [Gallid herpesvirus type 1] >gi 299459 bbs 129316 (S58431) CD4 precursor homolog to CD4 and IgM heavy chain [Gallid herpesvir hypothetical protein - Marek's disease virus gammaherpesvirus tumorigenicity associated mRNA, two complete cds's.], gene products [Gallid herpesvirus type 1] >gi 299459 bbs 129316 (S58431) CD4	0.93

SEQ ID	ACCESSION	DESCRIPTION1	P VALU
		STERYL-SULFATASE PRECURSOR (STEROID SULFATASE)	
656	1711563	(STERYL-SULFATE SULFOHYDROLASE) sulfatase [Mus	0.55
		HYPOTHETICAL 13.5 KD PROTEIN IN MOB1-SGA1	·
	A Memoran	INTERGENIC REGION >gi 626376 pir  S48473 probable membrane	***************************************
	мачи	protein YIL100w - yeast (Saccharomyces cerevisiae)	
657	731849	>gi 558707 emb CAA86281  (Z38125) orf, len: 117, CAI: 0.08, better	0.55
658	1669674	(X86819) Microtubule-associated protein 4	0.3
659	2493735	SKD3 PROTEIN SKD3 [Mus musculus]	0.25
660	110030	homeotic protein Hox 5.1 - mouse	0.19
TTTALLER NOTE THE CONTROL OF A STATE OF THE	**************************************	(AF000298) weak similarity to collagens; glycine- and proline-rich	/ · · · · · · · · · · · · · · · · · · ·
661	1947160	[Caenorhabditis elegans]	0.012
************	4567275	(AC006841) hypothetical protein [Arabidopsis thaliana]	0.005
*****		(AC002467) DRA protein (down-regulated in adenoma); sulfate	0.005
663	2677676	transporter; match to P40879 (PID:g729367) [Homo sapiens]	4e-056
	······································	hypothetical protein 2 - Mediterranean fruit fly	10 030
671	345474	; ·-	9.4
***************************************	4538951	(AL049488) putative protein [Arabidopsis thaliana]	9.1
	1749646	(D89219) unnamed protein product	7
~~~~	320302	ORF X protein - human papillomavirus type 41	5.5
·····		REPLICATION PROTEIN E1 papillomavirus (type 1)	J.J
675	267288	>gi 61013 emb CAA44657  (X62844) E1 [Pygmy chimpanzee	5.4
~~~	3581899	(AL031543) hypothetical serine-rich protein [Schizosaccharomyces	4.2
·····		(AL032653) predicted using Genefinder; cDNA EST EMBL:D36367	7.2
		comes from this gene; cDNA EST yk408c12.5 comes from this gene	
677	3881119		4.2
	***************************************	TRANSCRIPTIONAL ACTIVATOR FE65 APP interacting protein	<b>4.</b> 2
678	1169663	[Rattus rattus]	4
·	1565257	(X57108) cerebroside sulfate activator	3.2
~~~~	3093358	(AJ005559) SPR2A protein [Mus musculus]	1.1
interpretation of the contract	1763113	Commence and the commen	0.6
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3548791		0.46
·····	4567275		0.025
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3882311		7e-017
~~~~~~	2443342	on the contraction of the contra	5.1
		(D00232) E3 anti-[4-hydroxy-3-nitrophenyl(phenolate + phenolic	J.1
688	220464	2 32	6.4
•		HYPOTHETICAL PROTEIN MG181 Mycoplasma genitalium	U. <del>1</del>
689	1351502	A STATE OF THE STA	2.1
coccoccione and the second	250891	(32.2.2.2.)	4.6
		(AE001245) sugar ABC transporter, periplasmic binding protein	7.0
695	3323042		5.8
wa comercence with a series	1001741	an and a superior and	2.7
		DIHYDROXYACETONE KINASE (GLYCERONE KINASE)	L. 1
697	1169288	. !!	2~ 010
		DIHYDROXYACETONE KINASE (GLYCERONE KINASE)	3e-010
698	1169288	1400000 (7700004)	7- 010
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(Z49130) cDNA EST yk486b9.3 comes from this gene; cDNA EST	3e-010
699		140005	7- 006
**************************************			7e-005
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4504923	(AF007269) A IG002N01.1 gene product [Arabidopsis thaliana]	0.0004
	······································		1.5
	······	1	7.4
703.		(Z81531) F36D3.6 [Caenorhabditis elegans] HOMEOBOX PROTEIN HOX-D4 (HOX-4B) protein Hox D4 -	7.3
i			

SEQ ID		por (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VALU
Market arrange seminaran d	***************************************	DNA-directed RNA polymerase (EC 2.7.7.6) largest chain (isoform	
706	99899	C) - soybean (fragment) polymerase [Glycine max]	0.83
707	4263788	(AC006068) hypothetical protein	0.28
708	500858	(D14168) 50kDa lectin [Bombyx mori]	4e-011
***************************************	***************************************	(M30318) put. periplasmic receptor protein (chvE); putative	
711	142283	[Agrobacterium tumefaciens]	9.2
		collagen, type II, alpha 1 congenital)	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
		>gi 115287 sp P02458 CA12_HUMAN PROCOLLAGEN ALPHA	
712	4502949	1(II) CHAIN PRECURSOR [CONTAINS: CHONDROCALCIN] 1-	6.9
ONE SENSEMBLE CONTRACTOR OF THE PROPERTY OF	4539163	(AL049485) putative phytoene synthase	0.9
	1337103	(AC005990) Strong similarity to PFAM PF 00069 Eukaryotic protein	
715	4056437	kinase domain. [Arabidopsis thaliana]	
*********************	119296	ELASTIN PRECURSOR (TROPOELASTIN) gallus]	2.3
/ I U	119290		1.3
717	4226072	(AF125443) contains similarity to S. pombe phosphatidyl synthase	
/1/	4226073	(GB:Z28295) [Caenorhabditis elegans]	8e-016
		PEROXISOMAL MEMBRANE PROTEIN PMP30A protein - yeast	
710	2400761	(Candida boidinii) >gi 457391 (L27999) peroxisomal membrane	
/19	2498761	protein 31 [Candida boidinii]	1.3
720	2014062	SECRETORY CARRIER-ASSOCIATED MEMBRANE PROTEIN	
/20	3914963	3 >gi 2232237 (AF005036) secretory carrier membrane protein [Mus	0.58
701	4455004	(AF108226) immunoglobulin mu heavy chain precursor	***
	4457204	[Monodelphis domestica]	0.15
****	3860573	(AJ235270) unknown [Rickettsia prowazekii]	4.9
***************************************	2829912	(AC002291) Similar ATP-dependent RNA Helicase	0.0002
****	2130573	(U96771) putative polygalacturonase [Prevotella bryantii]	6.1
	539244	hypothetical protein YKR028w - yeast	6.1
	2633502	(Z99110) similar to hypothetical proteins from B. subtilis [Bacillus	4.7
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1078087	hypothetical protein YLR424w - yeast	1.6
729	4240219	(AB020672) KIAA0865 protein [Homo sapiens]	2
		(AB011874) alpha subunit of dinitrogenase reductase (Fe protein)	\$ }
***************************************	3165370	[unidentified nitrogen-fixing bacteria]	9.3
***************************************	3882195	(AB018280) KIAA0737 protein [Homo sapiens]	2e-061
xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx	3859938	(AF081101) reverse transcriptase [Lymantria dispar]	2.3
737	974143	(L42542) RLIP76 protein [Homo sapiens]	8.4
		(Z48583) similar to ATPases associated with various cellular	
		activities (AAA); cDNA EST EMBL:Z14623 comes from this gene;	
		cDNA EST EMBL:D75090 comes from this gene; cDNA EST	
	3877493	EMBL:D72255 comes from this gene; cDNA EST yk200e4.5	3e-047
739	4240235	(AB020680) KIAA0873 protein [Homo sapiens]	3e-052
740	4191810	(AB006532) DNA helicase [Homo sapiens]	1e-065
741	4507851	reserved protease [Homo sapiens]	2e-071
_		(Z81531) cDNA EST EMBL:D66579 comes from this gene; cDNA	**********
<b>3000</b>		EST EMBL:D70408 comes from this gene; cDNA EST yk263d3.5	
OLUMBA AND AND AND AND AND AND AND AND AND AN		comes from this gene; cDNA EST yk275c1.5 comes from this gene;	
742	3876797	cDNA EST EMBL:C10270 comes from this gene [Caenorh	2.7
ne verina esta esta esta esta esta esta esta est	4376875	(AE001642) CT465 hypothetical protein	0.53
······································	·······		0.32
		(U97016) similar to drosophila Rlc1 gene product ribosomal protein	U.J4
745		Y 4 /773 67 1) (2 775 )	0- 016
	iii ahaanaa aa a	***************************************	9e-016
·····			7.8
<del>~~~~</del>			7.8
, , ,		autotaxin precursor - human >gi 537906	6

EQ ID	ACCESSION	DESCRIPTION1	P VALU
		titin >gi 1212992 emb CAA62188  via Swiss-Prot; available at	
755	4507721	present via e-mail from LABEIT@EMBL-Heidelberg.DE [Homo	1.5
756	3036883	(AL022374) putative ABC transporter	0.68
757	543593	hypothetical 39.8K protein (clone GV-B) - garlic virus B	0.06
		hypothetical 30.5K protein - Enterococcus faecalis plasmid pAM-bet	a -
758	79960	1 >gi 3023044 (AF007787) orfC	1e-024
759	3882195	(AB018280) KIAA0737 protein [Homo sapiens]	6e-060
766	1707719	(Y08256) orf c02007 [Sulfolobus solfataricus]	9.8
767	2133808	immunoglobulin heavy chain - nurse shark	7.6
768	1469880	(D63483) The KIAA0149 gene product is related to Notch3. [Homo	7.5
769	4454062	(AJ132911) NorD protein [Bradyrhizobium japonicum]	5.9
		PROTEIN C2 >gi 74386 pir  WZVZB6 59K HindIII-C protein -	·
770	137532	vaccinia virus (strain WR)	4.4
Named and American Company	1098985	(U41031) proline-rich [Caenorhabditis elegans]	3.4
· · · · · · · · · · · · · · · · · · ·	4378891	(AF132481) Ese1L protein [Mus musculus]	2.6
·	**************************************	cone-rod homeobox PROTEIN >gi 2665534 (AF024711) cone rod	1
773	4557489	homeobox protein	1.5
·····	2135894	peripheral benzodiazepine receptor - human	0.52
museron man, 1949, Es	477495	cell-fate determining gene Notch2 protein - rat	0.51
		(U56965) Similar to NAD(P) transhydrogenase, mitochondrial; coded	
		for by C. elegans cDNA yk27c1.5; coded for by C. elegans cDNA	***
Other feet Walter		yk35b9.5; coded for by C. elegans cDNA yk35b9.3; coded for by C.	
776	1945493		10.20
······	746516	elegans cDNA yk161c9.3; coded for by C. elegans	0.39
///	740310	(U23517) D1022.7 [Caenorhabditis elegans] >gi 3258651 elegans]	0.059
		GLYCEROL-3-PHOSPHATE ACYLTRANSFERASE	
770	2400707	PRECURSOR (GPAT) (P90) acyltransferase homolog - mouse	
mmmmmmmmmm, n.j.,	2498786	>gi   193367 (M77003) glycerol-3-phosphate acyltransferase [Mus	0.058
	2134384	procKr2 - chicken (fragment) gallus]	0.015
and the second second second second	4176500	(AL031177) dJ889M15.3 (novel protein)	0.001
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	500858	(D14168) 50kDa lectin [Bombyx mori]	5e-010
······································	4204294	(AC003027) lcl prt_seq No definition line found	3e-010
·····	500858	(D14168) 50kDa lectin [Bombyx mori]	2e-010
morron con contrato,	3184082	(AL023781) N-terminal acetyltransferase 1	6e-014
·····	3184082	(AL023781) N-terminal acetyltransferase 1	3e-014
****	1330401	(U58762) T27F7.1 gene product [Caenorhabditis elegans]	6e-030
*************************	1330401	(U58762) T27F7.1 gene product [Caenorhabditis elegans]	3e-030
· · · · · · · · · · · · · · · · · · ·	3879850	(Z81592) predicted using Genefinder	8e-034
	1072198	(U40942) No definition line found [Caenorhabditis elegans]	1e-037
<del>~~~~~~</del>	3327160	(AB014573) KIAA0673 protein [Homo sapiens]	1e-062
······································	3413886	(AB007931) KIAA0462 protein [Homo sapiens]	1e-073
	1903264	(Y11824) hypothetical protein [Pisum sativum]	9.5
	***************************************	(AJ223176) Ser/Thr protein kinase	5.6
802	4521264	(AB015874) guanylate cyclase OIGC-R2	5.6
803 2	2144044	zinc finger protein AT-BP2 - black rat protein AT-BP2 [Rattus rattus]	
		(AF063866) ORF MSV092 hypothetical protein [Melanoplus	/
804	1049682	sanguinipes entomopoxvirus]	4.3
	***************************************	HYPOTHETICAL 73.3 KD PROTEIN C6G9.14 IN	
000	2842704	CHROMOSOME I >gi 1644326 emb CAB03616.1  pr otein	3.2
80512		0-1-1.10-1110-1110-111-11-11-11-11-11-11-11-11	ے. ب
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1526981		2.5
806 1	1526981 1082243	(X99945) amino acid permease YeeF like protein [Salmonella autotaxin precursor - human >gi 537906	2.5 2.5

SEQ ID	ACCESSION	or (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VALU
טויע זוט	ACCESSION	PHOSPHORIBOSYLAMINEGLYCINE LIGASE	IF VALU
	August - sager &	PHOSPHORIBOSYLFORMYLGLYCINAMIDINE CYCLO-	
	o Company	LIGASE (AIRS) synthetase, aminoimidazole ribonucleotide	*
	over a second	synthetase, glycinamide ribonucleotide transformylase {EC 6.3.4.13,	<u> </u>
800	2500002	į - · · · · · · · · · · · · · · · · · ·	2.4
009	2300002	6.3.3.1, 2.1.2.2} [Chironomus tentans, Peptide, 1371 aa titin >gi 1212992 emb CAA62188  via Swiss-Prot; available at	2.4
210	4507721	present via e-mail from LABEIT@EMBL-Heidelberg.DE [Homo	1.5
010	4507721	titin >gi 1212992 emb CAA62188  via Swiss-Prot; available at	1.5
011	4507721	present via e-mail from LABEIT@EMBL-Heidelberg.DE [Homo	1 4
011	430//21	INTERFERON-ACTIVATABLE PROTEIN 205 protein - mouse	1.4
		>gi 385703 bbs 133592 (S62227) D3=lipopolysaccharide-inducible	
812	2833215	[mice, macrophages, Peptide, 425 aa] [Mus sp.]	1.1
****	1262910	(U51645) cytidine triphosphate synthetase [Plasmodium falciparum]	
**********************	1245061	(U46069) fertilin alpha subunit [Oryctolagus cuniculus]	0.64
014	1243001	(AL034358) predicted using hexExon; L4830.10, Hypothetical	0.005
015	4493746		0.003
***************************************	3283350	protein, len: 816 aa [Leishmania major] (AF062378) calmodulin-binding protein SHA1 [Mus musculus]	
	4204294	omeronement commence and the commence of the c	0.003
******************		(AC003027) lcl prt_seq No definition line found	3e-006
reconnection and a series of the series of t	4309681 3293547	(AC006930) R33423_1 [Homo sapiens]	4e-007
***************************************		(AF072709) putative oxidoreductase [Streptomyces lividans]	5e-013
820	1086900	(U41278) contains similarity to G beta repeats (Z73428) similar to Zinc finger, C3HC4 type (RING finger); cDNA	2e-028
		EST EMBL:D67323 comes from this gene [Caenorhabditis elegans]	; ;
001	2070720	>gi 3881096 emb CAB16481  finger); cDNA EST EMBL:D67323	6 001
******************	3878739	comes from this gene	6e-031
	3882195	(AB018280) KIAA0737 protein [Homo sapiens]	1e-053
······	3095186	(AF057140) cargo selection protein TIP47 [Homo sapiens]	7e-060
	1915885	(Y08370) alpha-amylase [Crassostrea gigas]	9.2
842	1621461	(U73103) laccase [Liriodendron tulipifera]	9.2
942	222620	(S41487) possible ribosomal protein=I(3)S12 Canton S wild type	0.1
843	232620	[Drosophila melanogaster, Peptide, 73 aa] [Drosophila melanogaster] HYPOTHETICAL 73.6 KD PROTEIN CY49.21	9.1
044	1721220		
anno an anno anno an anno anno anno ann	1731338	>gi 1370248 emb CAA98194  (Z73966) hypothetical protein Rv2082	
843	3873261	(AF096295) cytochrome oxidase subunit I [Naja siamensis] (AF063866) ORF MSV173 putative serine/threonine protein kinase	6.9
		•	s \$
0.46	4040727	Swinepox virus C20L homolog (vaccinia F10L), similar to	
	4049727	SW:P32216 [Melanoplus sanguinipes entomopoxvirus]	5.5
	1175595	HYPOTHETICAL PROTEIN HI1265	4.1
848	446631	collagen:SUBUNIT=alpha2:ISOTYPE=IX [Homo sapiens]	4.1
0.40	(27171	odz protein - fruit fly (Drosophila sp.) product=tenascin homolog	
on our contraction of the contra	627171		3.2
850	3986440	(AF076785) serum amyloid A-activating factor SAF-5	3.1
0.51	01212	stem cell protein ERA-1-399, retinoic acid-induced - mouse	
······	91312	>gi 387146 (M22115) ERA-1-399 protein [Mus musculus]	1.4
:###.co.co.co.co.co.co.co.gc	4102043	(AF008203) homeobox protein [Homo sapiens]	1.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	746516	(U23517) D1022.7 [Caenorhabditis elegans] >gi 3258651 elegans]	0.28
	2935691	(AF032122) unknown [Streptococcus thermophilus bacteriophage	0.21
855		nodule-specific (hydroxy)proline-rich protein	0.014
~		DIHYDROXYACETONE KINASE (GLYCERONE KINASE)	
		>gi 493083 (U09771) dihydroxyacetone kinase	1e-005
••••••••••••••••••••••••••••••••••••••	***************************************	(AB017156) gob-5 [Mus musculus]	1e-011
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		(U29488) C56C10.3 gene product [Caenorhabditis elegans]	3e-012
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3293547	(AF072709) putative oxidoreductase [Streptomyces lividans]	5e-013
960	2228750	(U93868) RNA polymerase III subunit [Homo sapiens]	1e-015

SEQ ID	ACCESSION	DDDCKII ITOTTI	P VAL
	•	HYPOTHETICAL 24.2 KD PROTEIN C13A11.03 IN	
861	1175412	CHROMOSOME I >gi 984224 emb CAA90804	5e-021
		non-metastatic cells 5, protein expressed in (nucleoside-diphosphate	, , , , , , , , , , , , , , , , , , , ,
		kinase) KINASE HOMOLOG 5 (NDK-H 5) (NDP KINASE	*
883	4505413	HOMOLOG 5) nucleoside-diphosphate kinase [Homo sapiens]	9
New Action concerns the says and	3308984	(AB008516) mtprd [Mus musculus]	9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3273643	(AF042021) olfactory receptor [Sus scrofa]	
		HETEROCHROMATIN PROTEIN 1 HOMOLOG ALPHA (HP1	. <u> </u>
	name of the state	AI DUA) (ANTIGEN DOS) > -:1194211 (1.07515)	1
886	1170338	ALPHA) (ANTIGEN P25) >gi 184311 (L07515) complete cds.],	
880	1170336	gene product [Homo sapiens] chromosomal autoantigen [human,	9
		(U42832) coded for by C. elegans cDNA yk107c8.5; coded for by C	• *
007	1105745	elegans cDNA yk107c8.3; Similar to UDP-glucuronosyltransferase.	
60-00-00-00-00	1125745	[Caenorhabditis elegans]	6.9
	2921102	(AF020337) P6 [rice ragged stunt virus]	6.9
889	1707199	(U80841) C13A10.1 gene product [Caenorhabditis elegans]	6.8
		(U42832) coded for by C. elegans cDNA yk107c8.5; coded for by C.	
		elegans cDNA yk107c8.3; Similar to UDP-glucuronosyltransferase.	Parket Bayes
890	1125745	[Caenorhabditis elegans]	5.3
		PUTATIVE RIBOFLAVIN BIOSYNTHESIS ENZYME	
		>gi 1707704 emb CAA69508  (Y08256) riboflavin biosynthesis	
891	2493564	protein ribG [Sulfolobus solfataricus]	5.2
892	2276148	(Z81463) Similarity to C.elegans zinc finger proteins [Caenorhabditis	3.4 5.5.2
	**************************************	L-LACTATE DEHYDROGENASE 1.1.1.27) - Mycoplasma	5.7.2
893	1346425	genitalium (SGC3) >gi 1046180	5.0
	4234795	(AF078135) unknown [Leptospira borgpetersenii]	5.2
	·	(702780) cDNA EST EMDI D75052	5.2
895	3874963	(Z92780) cDNA EST EMBL:D75953 comes from this gene	***************************************
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3875723	[Caenorhabditis elegans]	4
·····		(Z54270) similar to membrane glycoprotein	3.9
orronover v v v manadajų.	1086593	(U41007) C33H5.15 gene product [Caenorhabditis elegans]	3.9
oraran meneraliya	3289979	(AC005263) SP62_HUMAN; SAP 62; SF3A66 [Homo sapiens]	3.1
·····	2208965	(Y10528) cyanide insensitive terminal oxidase [Pseudomonas	3
~~~~~~~~~~~	4539386	(AL035526) extensin-like protein	1.8
901	2662561	(AF036692) Similar to seven transmembrane receptor	1.8
AMORE		pheromone shutdown protein homolog - Methanococcus jannaschii	
- Children		>gi 1592009 (U67576) pheromone shutdown protein (traB)	
	2129184	[Methanococcus jannaschii]	1.3
	2317864	(U78289) tylactone synthase module 7 [Streptomyces fradiae]	0.61
	3808242	(AF069669) pol protein [Human immunodeficiency virus type 1]	0.6
	3327128	(AB014557) KIAA0657 protein [Homo sapiens]	0.46
	294529	(L14933) convertase PC5 [Rattus norvegicus]	0.35
907 3	3005087	(AF044924) hook2 protein [Homo sapiens]	0.12
908 3	3297817	(AL031032) putative protein [Arabidopsis thaliana]	0.025
-	and the second s	(AF043250) mitochondrial outer membrane protein [Homo sapiens]	0.023
ANAMAGA ATT	Reterior Company of the Company of t	>gi 3941347 (AF043253) mitochondrial outer membrane protein	•
909	941342	TTT	0.01
······································	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	finger protein rfp - mouse (fragment)	0.01
- 10	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		5e-006
0117	842526	(AL021746) hypothetical anaphase promoting factor component	
7112	·······	Schizosaccharomyces pombe] anaphase promoting complex	3e-018
012 4	226072	AF125443) contains similarity to S. pombe phosphatidyl synthase	
		GB:Z28295) [Caenorhabditis elegans]	4e-022
·····		AF058803) mucin 4 [Homo sapiens]	3e-038
914 3	882195 (	AB018280) KIAA0737 protein [Homo sapiens]	4e-051
·····			
	[]	RNA-directed RNA polymerase (EC 2.7.7.48) - Marburg virus (strain Musoke) >gi 332179 (M92834) L Protein [Marburg virus]	

SEQ ID	ACCESSION	DESCRIPTION1	P VALU
		CYTOCHROME B >gi 2144288 pir  H22848 ubiquinolcytochrome-	
941	117865	c reductase (EC 1.10.2.2) cytochrome b - Sauroleishmania tarentolae	8.6
1		(AL008970) predicted using hexExon; MAL3P4.10 (PFC0515c),	,
942	3764008	Hypothetical protein, len: 1237 aa	8.6
	**************************************	(Z98547) predicted using hexExon; MAL3P3.11 (PFC0380w), Dual-	ф
ĺ		specificity protein phosphatase, len: 581 aa; Similarity to protein	, Waterparen
		phosphatases. S.cerevisiae protein-tyrosine phosphatase YVH1	T TOTAL CONTROL OF THE CONTROL OF TH
943	3649770	(SW:PVH1 YEAST) BLAST Score: 123, s	6.5
		(U42832) coded for by C. elegans cDNA yk107c8.5; coded for by C.	
		elegans cDNA yk107c8.3; Similar to UDP-glucuronosyltransferase.	
944	1125745	[Caenorhabditis elegans]	5.1
***************************************	2208965	(Y10528) cyanide insensitive terminal oxidase [Pseudomonas	5
	2146997	L-selectin precursor - rabbit >gi 847788	,5
ノサリ	2140771	ALANYL-TRNA SYNTHETASE (ALANINETRNA LIGASE)	
7		(ALARS) >gi 95227 pir  S16897 alaninetRNA ligase (EC 6.1.1.7) -	***
047	125001		2.0
94 /	135091	Rhizobium leguminosarum bv. viciae synthetase [Rhizobium	3.8
		NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4	
		>gi 102585 pir  S26021 NADH dehydrogenase mitochondrion	
948	1171815	(SGC4) >gi 559495 emb CAA38170  (X54253) ND4 protein [Ascaris	3.8
10000		glycoprotein B homolog precursor - feline herpesvirus 1	and a second
		>gi 261095 bbs 120003 (S49775) glycoprotein B homolog [feline	
949	2120034	herpesvirus type 1 FHV-1, Peptide, 948 aa] [Feline herpesvirus 1]	1.7
demonstrate		UNKNOWN >gi 2587054 (AF027204) putative tetraspan	9
950	4507539	transmembrane protein L6H [Homo sapiens]	0.99
Www.		CHEMOTAXIS LAFT PROTEIN parahaemolyticus >gi 677909	
distribution of the second	417237	(U20541) LafT [Vibrio parahaemolyticus] >gi 1518953 (U52957)	0.26
952	3798624	(AF082100) FK506 polyketide synthase [Streptomyces sp. MA6548]	0.12
		(AL034358) predicted using hexExon; L4830.10, Hypothetical	
953	4493746	de Carrent de la companyament de	0.003
S. West o		HYPOTHETICAL 24.2 KD PROTEIN C13A11.03 IN	
*************************************	1175412	CHROMOSOME I >gi 984224 emb CAA90804	6e-014
955	4240235		8e-044
1 1		RIBONUCLEOSIDE-DIPHOSPHATE REDUCTASE ALPHA	
disease for.		CHAIN (RIBONUCLEOTIDE REDUCTASE) reductase (EC	
976	1350599	1.17.4.1) - Mycoplasma genitalium (SGC3) reductase, alpha chain	8.2
Miles and a		HYPOTHETICAL 23.9 KD PROTEIN IN SGA1-KTR7	
MMI, Jane		INTERGENIC REGION >gi 1077785 pir  S49791 probable	
· · · · · · · · · · · · · · · · · · ·		membrane protein YIL089w - yeast (Saccharomyces cerevisiae)	
977	731840	>gi 577125 emb CAA86705.1  (Z46728) YI9910.07, unknown orf,	6.4
978	2132436	probable membrane protein YDL118w - yeast	6.3
979	1710518	60S RIBOSOMAL PROTEIN L22 homologue to human L22	6.3
980	3602956	(AF041468) rbcR homolog [Guillardia theta]	4.9
	2146997	L-selectin precursor - rabbit >gi 847788	4.8
***************************************	ts e en el 18 le en en el la librar en el la latera en el la en	(AL023838) predicted using Genefinder; similar to Helix-hairpin-	~~
to Visitana-Po		helix motif.; cDNA EST yk241d12.5 comes from this gene; cDNA	
a qualitive		EST yk401c3.5 comes from this gene; cDNA EST CEMSE47F	
982	3880890	comes from this gene [Caenorhabditis elegans]	4.8
Witness and Witness and Park agency and pro-	3880727	(AL032632) predicted using Genefinder	2.8
983		NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2	
983			
983		>gi 2147757 njr  S62766 NADH dehydrogenase caldarium)	
	1352547	>gi 2147757 pir  S62766 NADH dehydrogenase caldarium)	2.2
984	1352547	mitochondrion (fragment) subunit 2 [Cyanidium caldarium]	2.2
984	1352547 3551845		2.2

mon-witomonwood	ACCESSION	DESCRIPTION1	P VAL
harfulge the next presuppress out the	1707126	(U80454) T16A1.2 [Caenorhabditis elegans]	2.1
****	1497971	(U55797) VP6 [Bluetongue virus]	1.7
	3877568	(Z70208) similar to collagen	0.97
990	3309543	(AF036382) MLL [Fugu rubripes]	1e-005
991	4240235	(AB020680) KIAA0873 protein [Homo sapiens]	2e-039
1015	3257605	(AP000005) 484aa long hypothetical protein	8.2
	na-	(X57019) unnamed protein product [Homo sapiens]	
	***************************************	>gi 238775 bbs 65126 (S65125) putative tyrosine kinase	
1016	37593	receptor=UFO [human, NIH3T3, Peptide, 894 aa]	8.1
1017	746533	(U23520) similar to cuticular collagen [Caenorhabditis elegans]	8
1018	729045	BETA CASEIN PRECURSOR scrofa]	7.9
		INTERLEUKIN-1 BETA CONVERTASE PRECURSOR (IL-1BC)	
1019	1170463	(IL-1 BETA CONVERTING ENZYME) (ICE) converting enzyme	7.8
1020	1197641	(U46859) DdhB [Yersinia enterocolitica (type 0:8)]	6
1021	2104691	(U92794) alpha glucosidase II, beta subunit [Mus musculus]	4.7
1022	1139577	(D63706) Orf5 [Streptomyces griseus]	4.7
		EARLY GLYCOPROTEIN GP48 PRECURSOR glycoprotein (18)	
1023	136785	[human herpesvirus 5]	3.5
1024	1314757	(U54761) phosphoglucose isomerase [Erwinia amylovora]	2.6
***************************************		PUTATIVE TRANSPOSASE FOR INSERTION SEQUENCE IS408	
		>gi 309869 (L09108) IS408 transposase; putative [Pseudomonas	
1025	2497383	cepacia] >gi 1097384 prf  2113421C transposase [Burkholderia	1.6
		CHITOOLIGOSACCHARIDOLYTIC BETA-N-	······
		ACETYLGLUCOSAMINIDASE PRECURSOR (BETA-	
		GLCNACASE) 3.2.1) - silkworm >gi 998377 bbs 165703 (S77548)	
1026	1346281	chitooligosaccharidolytic beta-N-acetylglucosaminidase, beta-	0.7
		ALPHA-1D ADRENERGIC RECEPTOR (ALPHA 1D-	
		ADRENOCEPTOR) (ALPHA-1A ADRENERGIC RECEPTOR)	
1027	3121722	receptor [mice, brain, Peptide, 562 aa] [Mus sp.]	0.18
1028	4071321	(AF073954) Y-box protein MSY2 [Mus musculus]	0.14
1029	1293835	(U56965) C15H9.5 gene product [Caenorhabditis elegans]	0.11
	***************************************	(AC004084) similar to GTPase-activating proteins; 35% similar to	***************************************
1030	2822157		2e-013
1040	1877301	(Z92774) hypothetical protein Rv3570c	9.7
		GAMMA-SOLUBLE NSF ATTACHMENT PROTEIN (SNAP-	Medicine del processor conservago appaya, es
TA PAGE		GAMMA) >gi 423252 pir  S32369 gamma-SNAP protein - bovine	
and the same		>gi 298669 bbs 127528 gamma soluble NSF attachment protein,	
1041	3024629	gamma SNAP=N-ethyl-maleimide-sensitive fusion protein attachment	7.8
		UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 7	************************
		(UBIQUITIN THIOLESTERASE 7) probable membrane protein	
		YIL156w - yeast (Saccharomyces cerevisiae)	
1042	731044	>gi 557767 emb CAA86122  (Z38059) orf, len: 1071, CAI: 0.13,	7.6
1043	2190592	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4.3
1044	2627435	and the contract of the contra	3.5
1045	3881166	<del>งาวอาการและสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามารถสามา</del>	2
		LA PROTEIN HOMOLOG (LA RIBONUCLEOPROTEIN) (LA	**************************************
1046	729918	· · · · · · · · · · · · · · · · · · ·	0.009
	······································	(Z83819) dJ146H21.2 (similar to CYTOCHROME B-245 HEAVY	
1047	4106562		3e-017
	***************************************	down-regulated in adenoma protein down-regulated in adenoma	
diagnostiva		(DRA) - human >gi 291964 576-580, 579-583; acidic transcr. activ.	
1048	4557535		4e-026
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4240235		1e-020
1049	4240233	(ADV2VVVV) KIAAVV/ 3 DIVICIII [1]VIIIO SADIEIISI	

EQ ID	ACCESSION	bor (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VAL
	THE PARTY OF THE P	VITAMIN K-DEPENDENT GAMMA-CARBOXYLASE (GAMMA	Tr AVT
	(April 1970)	GLUTAMYL CARBOXYLASE) 4.1.1) - bovine >gi 289399	•
1065	586240	(L09726) gamma-carboxylase	0.6
******************	3413410	(AL031231) 30S ribosomal protein S15	9.6
~~~~~~~~~		GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2	7.4
		ANTIGEN) (IGE-BINDING PROTEIN) (35 KD LECTIN)	Avenue and a second
1067	1170758	(CARROLLIND ATE DARRING PROTEIN) (35 KD LECTIN)	about the same of
		(CARBOHYDRATE BINDING PROTEIN 35) (CBP 35)	7.4
1008	867985	(L39002) S1 gene product [Avian orthoreovirus] orthoreovirus]	7.4
1060	107404	METALLOTHIONEIN (MT) lucius] >gi 62783 emb CAA49636	
1069	127494	(X70042) Metallothioein	7.3
		(AL023634) palmitoyl-protein thioesterase precursor	
	3150259	[Schizosaccharomyces pombe]	5.7
1071	328647	(M81729) tat [Human immunodeficiency virus type 1]	4.3
1072	2605979	(AF030027) 35 [Equine herpesvirus 4]	3.3
		PROTEIN C2 >gi 74386 pir  WZVZB6 59K HindIII-C protein -	<b></b>
1073	137532	vaccinia virus (strain WR)	3.3
1074	3769620	(AF091563) olfactory receptor [Rattus norvegicus]	3.3
***************************************	*******************************	(U15829) envelope glycoprotein V3-V5 loop region [Human	3.3
1075	687367	immunodeficiency virus type 1]	
		TRANSCRIPTION INITIATION FACTOR TFIID 60 KD	1.4
1076	3915901	SUBUNIT (TAFII-60) (TAFII60)	
	2317934	(IAFII-00) (IAFII00)	0.48
	3878966	(U97553) unknown [murine herpesvirus 68]	0.37
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	***************************************	(Z68009) R09A8.1 [Caenorhabditis elegans]	9.1
	2341037	(AC000104) F19P19.17 [Arabidopsis thaliana]	2.3
1084	4508019	zinc finger protein 231 protein [Homo sapiens]	8.9
The second		(Z69903) predicted using Genefinder; Similarity to Rat casein kinase	<b>70-40</b>
i i i i i i i i i i i i i i i i i i i		I (SW:KC1D_RAT); cDNA EST EMBL:D65322 comes from this	
Management		gene; cDNA EST EMBL:D68704 comes from this gene; cDNA EST	
1085	3877198	yk475f2.5 comes from this gene [Caenorhabditis	4
and the same of th		(J02691) mitochondrial phenylalanyl-tRNA synthetase alpha subunit	*
1086	171998	precursor [Saccharomyces cerevisiae]	2.3
		LARGE TEGUMENT PROTEIN BPLF1 reading frame, 1 NXT/S,	
1087	135574	TITLE DECEMBER OF THE FEE	0.77
1091	1718312	/T TO C ( O O O O O T T T O O O O O O O O O O O	9.8
~~~~	······································	stress-sensitive restriction system protein 2 - Corynebacterium	9.0
WWwww		glutamicum (ATCC 13032) >gi 549844 restriction endonuclease	
1092	1076012	vehicle is store	
	882341	(II) (1700) I D D 1 FA 1 · 1 · 1 · 1	7.5
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		( A TOO 5 43 0 ) 1	0.71
*************************	·····	(AJ005438) beta adrenoreceptor B	0.54
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	La constitución de la constituci	(U41278) contains similarity to G beta repeats	1e-028
	4519908	(AB018106) HrSmad1/5 [Halocynthia roretzi]	6
and the second second second second	3043868		5.9
1098 2		NEUROGENIC LOCUS NOTCH 3 PROTEIN	3.5
Villa villa and	, , , , , , , , , , , , , , , , , , ,	(Z72514) predicted using Genefinder; similar to collagen; cDNA	
	MMMoonm	EST EMBL:M89258 comes from this gene; cDNA EST	
***************************************	W. scooning	EMBL:D68856 comes from this gene; cDNA EST yk232e11.3 comes	
1099 3	8879748	from this case. DNA Figm 1000 44 5	0.079
1100 1	123087	(U42436) C49H3.3 gene product [Caenorhabditis elegans]	5e-006
	**	(AL021483) similar to Probable rabGAP domains; cDNA EST	~~~~~~
о памене	very sono	EMBL:D34945 comes from this gene; cDNA EST EMBL:D27313	
	· ·	comes from this gene; cDNA EST EMBL:D2/313	
1101 3	880809	TALL FOR DAMP PORCE	
~~~~~~~~~~~	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1170664) 2 keto 2 doowych again 11 m ffy 1 6	Se-011
точностичностью общения	aramacan recommenda recommendad	A CTINI DIDITO DE CATERIO	5.8
LIOSEL			5.5

	ACCESSION	or (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VALU
	2702276	(AC003033) putative beta-glucosidase [Arabidopsis thaliana]	4.2
announcement and a second and a	1504006	(D86966) similarto human ZFY protein. [Homo sapiens]	4.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2337833	(Z98604) hypothetical protein MLCB2052.27	1.8
	3122952	TIPD PROTEIN >gi 2407788 (AF019236) TipD [Dictyostelium]	6e-019
	1049344	(AL034567) putative protein [Arabidopsis thaliana]	6.8
	**************************************	glycoprotein - rat >gi 986943 (L08134) glycoprotein [Rattus	
1113 2	2143767	norvegicus] norvegicus]	0.018
1116 2	THE TAXABLE PROPERTY OF THE PARTY OF THE PAR	(M63849) 2'-5' oligoadenylate synthetase [Mus musculus]	7.9
1119 1		MITOCHONDRIAL RIBOSOMAL PROTEIN VAR1 1-339)	9.7
rancom war aware as according	580768	(X92688) NADH dehydrogenase [Pecten maximus]	4.4
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	587868	(AF069302) Orf2 [Pediococcus pentosaceus]	1.5
1121		(AF051933) cyclin T; positive elongation transcription factor b cyclin	
1123 3	851496	subunit [Drosophila melanogaster]	8
**************************************	877133	(Z83109) predicted using Genefinder	3.4
1131 3	1077133	DNA POLYMERASE (A PROTEIN) 2.7.7.7) - ground squirrel	6
1132 1	19964	hepatitis virus >gi 325402	1
1133 4	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		4.6
named a recommence of the configuration of the conf	100099	(U03388) cyclooxygenase 1 [Rattus rattus]	4.5
<del>~~~~~~</del>	130154	(U93502) hypothetical protein 256 [Secale cereale]	2.7
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	~~~~~	hypothetical protein 241 - maize chloroplast	1.2
	521280	(AB011832) cytochrome c oxidase subunit I [Dicyema misakiense]	0.94
******	706875	(D85084) NCAM-180 [Cynops pyrrhogaster]	0.68
1138 7	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(U23517) D1022.7 [Caenorhabditis elegans] >gi 3258651 elegans]	0.061
1139 7	46516	(U23517) D1022.7 [Caenorhabditis elegans] >gi 3258651 elegans]	0.061
		(U97016) similar to drosophila Rlc1 gene product ribosomal protein	
1140 1	938549	L4 (YML4) (NID:g459259)	5e-017
		(U97016) similar to drosophila Rlc1 gene product ribosomal protein	
	······································	L4 (YML4) (NID:g459259)	3e-017
anno anno anno anno anno anno anno anno		(AF100421) p80 [Rattus norvegicus]	1e-056
<del>vana vano en </del>		(AB006532) DNA helicase [Homo sapiens]	8e-066
		(AF100421) p80 [Rattus norvegicus]	8e-070
1163 82		probable serine-type carboxypeptidase (EC 3.4.16.1) - wheat	9.7
1164 3	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(Z69885) predicted using Genefinder	9.6
1165 10	·**	mercury resistance regulatory protein KW20)	9.5
			9.3
1167 48		(L20303) actin filament-associated protein [Gallus gallus]	7.6
		trypsin inhibitor, WTI [Triticum aestivum=wheat, variety San	
1168 50		Pastore, endosperm, Peptide, 71 aa]	7.5
***************************************		(AE001240) spermidine/putrescine ABC transporter, permease	
1169 33		protein (potB) [Treponema pallidum]	7.5
1170 11	707179	(U80839) ZC204.15 gene product [Caenorhabditis elegans]	7.4
1171 20	09619	(J01901) major coat protein A [Adeno associated virus 2]	7.3
1172 16		(D88529) serine acetyltransferase serine acetyltransferase [Spinacia	7.3
1173 72	22375	(U23139) similar to phospholipase ADRAB-B precursor	7.2
	1	TAIL FIBER PROTEIN GP37 (RECEPTOR RECOGNIZING	***************************************
000	i si contra	PROTEIN) >gi 76090 pir  TLBP74 tail fiber protein gp37 - phage T4	
		>gi 15374 emb CAA24228  (V00863) gene 37 [coliphage T4]	
1174 13			5.5
1175 10		(D64003) hypothetical protein	4.4
••••••••••••••••••••••••••••••••••••••	····	HYPOTHETICAL 126.5 KD PROTEIN C31A2.16 IN	······································
1176 11			4.4
1177 28	329216	(AF044287) delta adaptin subunit of AP-3 [Drosophila melanogaster]	
		histone-lysine N-methyltransferase (EC 2.1.1.43) large chain N-	7.4
***************************************	, , , , , , , , , , , , , , , , , , ,	methyltransferase - garden pea carboxylase large subunit N-	
1178 10			4.1
11/0/10	/UT-//T	menymanoterase [r tsum sanvum]	4.1

Corresp   feature   HYPO   CHRO   (U404   elegans   55   repeats   53   (AF06)   (U404)   elegans   55   repeats   61   (AF118)	DESCRIPTION1 72) This CDS feature is included to show the translation of the ponding V_region. Presently translation qualifiers on V_region are illegal. THETICAL 49.3 KD PROTEIN C09G5.1 IN MOSOME II >gi 3874104 emb CAA86760  10) coded for by C. elegans cDNA yk9e10.5; coded for by C. as cDNA yk9e10.3; multiple regions of similarity to EGF-like and cysteine-rich repeats 8182) B cell linker protein BLNK [Mus musculus] 10) coded for by C. elegans cDNA yk9e10.5; coded for by C.	3.8 3.2 2.6 2.6
Corresp   feature   HYPO   CHRO   (U404   elegans   55   repeats   53   (AF06)   (U404)   elegans   55   repeats   61   (AF118)	ponding V_region. Presently translation qualifiers on V_region is are illegal.  THETICAL 49.3 KD PROTEIN C09G5.1 IN  MOSOME II >gi 3874104 emb CAA86760   10) coded for by C. elegans cDNA yk9e10.5; coded for by C. is cDNA yk9e10.3; multiple regions of similarity to EGF-like is and cysteine-rich repeats  8182) B cell linker protein BLNK [Mus musculus]	3.8
6 feature	es are illegal.  THETICAL 49.3 KD PROTEIN C09G5.1 IN  MOSOME II >gi 3874104 emb CAA86760   10) coded for by C. elegans cDNA yk9e10.5; coded for by C. s cDNA yk9e10.3; multiple regions of similarity to EGF-like and cysteine-rich repeats  8182) B cell linker protein BLNK [Mus musculus]	3.8
HYPO CHRO (U404 elegans repeats (U404) elegans repeats (U404) elegans repeats (AF118)	THETICAL 49.3 KD PROTEIN C09G5.1 IN MOSOME II >gi 3874104 emb CAA86760  10) coded for by C. elegans cDNA yk9e10.5; coded for by C. s cDNA yk9e10.3; multiple regions of similarity to EGF-like and cysteine-rich repeats 8182) B cell linker protein BLNK [Mus musculus]	3.2
25 CHRO (U404 elegans 55 repeats (U404 elegans (U404 elegans repeats (AF118	MOSOME II >gi 3874104 emb CAA86760  10) coded for by C. elegans cDNA yk9e10.5; coded for by C. s cDNA yk9e10.3; multiple regions of similarity to EGF-like and cysteine-rich repeats 8182) B cell linker protein BLNK [Mus musculus]	2.6
(U404 elegans 55 repeats (U404 elegans 55 repeats 61 (AF118	10) coded for by C. elegans cDNA yk9e10.5; coded for by C. s cDNA yk9e10.3; multiple regions of similarity to EGF-like and cysteine-rich repeats  8182) B cell linker protein BLNK [Mus musculus]	2.6
elegans 55 repeats 53 (AF06) (U404) elegans 55 repeats 61 (AF118)	s cDNA yk9e10.3; multiple regions of similarity to EGF-like and cysteine-rich repeats 8182) B cell linker protein BLNK [Mus musculus]	~~~
55 repeats 53 (AF06) (U404) elegans 55 repeats 61 (AF118)	s and cysteine-rich repeats 8182) B cell linker protein BLNK [Mus musculus]	~~~
53 (AF06) (U404) elegans 55 repeats 61 (AF118)	8182) B cell linker protein BLNK [Mus musculus]	~~~
(U404) elegans repeats 61 (AF118)	10) coded for by C. elegans cDNA vk9e10.5; coded for by C.	
elegans 55 repeats 61 (AF118	to coded for by c. elegans china yksetu.s. coded for by (	<b></b>
55 repeats 61 (AF118	s cDNA yk9e10.3; multiple regions of similarity to EGF-like	LL CANADA
61 (AF118	and cysteine-rich repeats	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3391) salivary peroxidase	2.5
9 mating	5391) salivary peroxidase	2.5
	-type locus protein b3 - smut fungus	2.1
	84) polyprotein [Hepatitis C virus]	<u>‡</u> 1.5
~~~~	3844) similar to Ion transport proteins [Caenorhabditis	1.4
21 (AF053	3091) eyelid [Drosophila melanogaster]	1.4
commence and the second se	(2773) basic helix-loop-helix transcription factor Ndr1b [Danio	1.4
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		0.62
	2) cutinase negative acting protein [Fusarium solani f. sp.	0.5
(Z/020	7) predicted using Genefinder; similar to collagen; cDNA	
EST EN	MBL:D65905 comes from this gene; cDNA EST	
EMBL:	D65858 comes from this gene; cDNA EST EMBL:D69306	
)4 comes t	from this gene; cDNA EST EMBL:D65755 comes from this	0.025
(AF067	(217) contains similarity to myosin head (motor) domains	\$
(Pfam:	myosin_head.hmm score: 602.72, 40.38 and 128.290) and	
phorbol	esters/diacylglycerol binding domains (Pfam: DAG PE-	; 1
		0.002
***************************************	rich protein - mouse proline-rich protein [Mus musculus]	5e-005
7 (AF049	910) TACC1 [Homo sapiens]	7e-011
(AF014	903) NADH dehydrogenase subunit 2 [Pan troglodytes]	
6 >gi 328	7338 (AF014904) NADH dehydrogenase subunit 2 [Pan	5e-011
(AF125	443) contains similarity to S. pombe phosphatidyl synthase	}
3 (GB:Z2	8295) [Caenorhabditis elegans]	2e-011
(U2948	8) C56C10.3 gene product [Caenorhabditis elegans]	2e-011
3 (AB020	063) Keap1 [Mus musculus]	5e-011
PUTAT	IVÉ AMÍNOPEPTIDASE ZK353.6 IN CHROMOSOME III	30 012
>gi 1078	8908 pir S44657 ZK353.6 protein - Caenorhabditis elegans	1
>gi 289	760 (L15313) homology with lengthe aminopentidase; coded	
for by C	elegans cDNAs CE2F12 (GenBank: 714714) and CE15D11	20 014
	768) brain specific cortactin-hinding protein CRP90 (Pottus	2e-014
	709) putative exidereductase [Streptomyces lividage]	2e-016 2e-019
······································	568) tumor suppressing STF cDNA 4 [Home senions]	*****
·····	>gi 3928762 dhi BAA34703	3e-031
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		1e-048
	0000111 10001 100001 100001	4 071
······································	4.0.4.\	4e-051
		2e-064
(ABUI /)	) putative Inositol polyphosphate phosphatase	3e-071
	pulative illustroi polyphosphate phosphatase	
(Z98/03	•	
S [Schizos	accharomyces pombe]	9.1
S [Schizos NADH o	accharomyces pombe] dehydrogenase subunit I - maize	9.1 8.3 8.3
	50 (AF115) 2 (M132) 51 (U6184) (Z7020) EST EN EMBL: 04 comes 1 (AF067) (Pfam: phorbol) bind.hn proline- 57 (AF049) (AF014) 56 >gi 328 (AF125) (3 (AB020) PUTAT >gi 1073 >gi 2897 for by C 2 (AF053) 7 (AF072' 8 (AF125) 1 reserved	(AF115773) basic helix-loop-helix transcription factor Ndr1b [Danio (M13235) abl-like putative oncogene; putative (U61842) cutinase negative acting protein [Fusarium solani f. sp. (Z70207) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D65905 comes from this gene; cDNA EST EMBL:D65858 comes from this gene; cDNA EST EMBL:D69306 comes from this gene; cDNA 4 [Homo sapiens]

		oor (BlastX vs. Non-Redundant Proteins)	
SEQ ID	ACCESSION		P VALU
		C4B-BINDING PROTEIN BETA CHAIN PRECURSOR	The formation of the first section of the section o
	and the definition of the second	>gi 2143627 pir  S57960 C4BP protein beta chain precursor - rat	- Andrews
	2493794	>gi 899382 emb CAA90392  (Z50052) C4BP beta chain, leader	8.2
1250	1703342	APOLIPOPROTEIN D PRECURSOR	6.8
		PROBABLE RNA-DIRECTED RNA POLYMERASE polymerase	***************************************
1251	133652	(EC 2.7.7.48) - southern bean mosaic virus mosaic virus]	6.6
		hypothetical protein MJ0749 - Methanococcus jannaschii	***************************************
1252	2128472	>gi 1591462 (U67520) conserved hypothetical protein	6.5
		NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4	
1253	2499265	>gi 1182022 emb CAA56534  (X80245) ND4 [Locusta migratoria]	6.4
1254	1098322	Ran/TC4-binding nucleopore protein [Homo sapiens]	5.3
		homeotic protein smox-5 - fluke (Schistosoma mansoni) >gi 161110	
1255	283518	(M85304) homeodomain protein	5.2
		EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-	
		RICH GLYCOPROTEIN) >gi 82169 pir  S06733 hydroxyproline-rich	1
9		glycoprotein precursor - common tobacco >gi 19867 emb CAA32090	-; [[
1256	119714	(X13885) extensin (AA 1-620) [Nicotiana tabacum]	4.9
·····	and the second contract of the second contrac	HYPOTHETICAL 100.9 KD PROTEIN C34E10.3 IN	7.7
1257	1176529	CHROMOSOME III >gi 500726 (U10402) C34E10.3 gene product	4.8
	***************************************	COPPER RESISTANCE PROTEIN B PRECURSOR	<u> </u>
1258	116922	>gi 77826 pir  B32018 copper resistance protein B precursor -	3.8
	**************************************	HYPOTHETICAL 19.0 KD PROTEIN IN NNF1-STE24	3.0
hubolandene		INTERGENIC REGION >gi 1077909 pir  S57138 hypothetical	
designa		protein YJR115w - yeast (Saccharomyces cerevisiae)	
1259	1352916	>gi 1015833 emb CAA89645  (Z49615) ORF YJR115w	3.7
·	184111	(M20677) Kruppel-related protein (AA at 172) [Homo sapiens]	3.7
	***************************************	BOLA CLASS I HISTOCOMPATIBILITY ANTIGEN, ALPHA	3.1
1261	122137	CHAIN BL3-6 PRECURSOR antigen alpha chain precursor (BL3-6)	2.6
~~~~		(U41263) this gene lies in the intron of T19D12.4 and on the opposite	73.0
		strand; strong similarity to casein kinases and to C. elegans proteins	Block of
1262	1086822	C03C10.2, F41G3.5 and ZK507.1	2
····		(AP000003) 338aa long hypothetical protein	3
		(AL034558) predicted using hexExon; MAL3P2.18 (PFC0245c),	3
1264		Hypothetical protein, len: 3934 aa	20
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	GENERAL STRESS PROTEIN A Bacillus subtilis	2.8
1265		>gi 580866 emb CAA51568  (X73124) ipa-12d [Bacillus subtilis]	2.2
	/ <del>////////////////////////////////////</del>		2.2
immercano constitución de la const		(AB001684) ORF54c [Chlorella vulgaris]	2.1
	<del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	(AF079369) transcriptional repressor TUP1 [Dictyostelium	1.4
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(A FOCOGOC) DOANS	1.4
	and construction and a second	/7.1.(A=A) () (A=A) (A=A)	1.3
		(ATO702 (A)	1.3
	**************************************		1.2
1272	2145956	probable phosphomannomutase (EC 5.4.2.8) - Mycobacterium leprae	•
annan anna ann ann ann ann ann ann ann	and a second and a	>gi 467178 (U00022) u0308b; L308 F1_13 [Mycobacterium leprae] (AE001012) GMP synthase (guaA-2) [Archaeoglobus fulgidus]	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
reconstruction of the same and the same		( ) = 0.0 ( ) = - \	1
······································	<del></del>	(200010)	0.76
······································	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		0.21
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(T T 4 A 4 A C ) C 4 A T T A A	0.14
		T T A O A O C \ O A O T T O A	0.013
1279 1	<del>~~~</del>	3 - ( 0 0 - 1 )	0.0004
***************************************	······································	TO FA	7e-005
***************************************	608372	1 700 700 10 77 77 77	9e-011
<del>maraninina makalangan di</del> kaca	6114818 (	AJ005891) JM2 [Homo sapiens]	5e-011
1282 3	608372 (	AF053768) brain specific cortactin-binding protein CBP90 [Rattus	7e-016

	ACCESSION	por (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VALI
1283	3608372	(AF053768) brain specific cortactin-binding protein CBP90 [Rattus	2e-016
1284	3649741	(AJ000281) mucin [Homo sapiens]	2e-031
1285	4240307	(AB020716) KIAA0909 protein [Homo sapiens]	1e-043
1286	4240307	(AB020716) KIAA0909 protein [Homo sapiens]	1e-043
1287	3649741	(AJ000281) mucin [Homo sapiens]	1e-047
······	N. Marie Carlo Car	down-regulated in adenoma protein down-regulated in adenoma	16-04/
THE STATE OF THE S		(DRA) - human >gi 291964 576-580, 579-583; acidic transcr. activ.	
1288	4557535	domain 620-640,; homeobox motif 653-676 [Homo sapiens]	le-060
		(Z50875) Proline rich domain; cDNA EST EMBL:D35637 comes	16-000
		from this gene; cDNA EST yk322a3.5 comes from this gene; cDNA	S S-
		EST yk397b2.5 comes from this gene; cDNA EST yk348b11.5	ann in
1324	3879589	comes from this gene; cDNA EST yk397b2.3 comes from	0.0
www.	3879505	(AL023816) T05G11.4 [Caenorhabditis elegans]	9.9
·	1346035	FOS-RELATED ANTIGEN-1 >gi 998348	9.4
1520	1340033	(AE000981) dipeptide ABC transporter, dipeptide-binding protein	8
1327	2648784		
	142774	[(dppA) [Archaeoglobus fulgidus]	7.9
1320	142//4 **** *******************************	(L07023) delta-endotoxin [Bacillus thuringiensis]	7.9
1220 2	2648784	(AE000981) dipeptide ABC transporter, dipeptide-binding protein	to to tolerana
~~~~~~~~ <u>~~~</u>		(dppA) [Archaeoglobus fulgidus]	7.9
·····	3874201	(Z81015) predicted using Genefinder	7.7
emorana waxaa aa aa aa gara	2358287	(AF010404) ALR [Homo sapiens]	7.7
····	3881262	(AL021175) Y6E2A.6 [Caenorhabditis elegans]	6
marrows and a second proper	974143	(L42542) RLIP76 protein [Homo sapiens]	6
1334 3	3287188	(Y10601) ankyrin-like protein [Homo sapiens]	6
100-		UBIQUITINPROTEIN LIGASE RSP5 yeast (Saccharomyces	
1335 7	30684	cerevisiae) >gi 603364 (U18916) Rsp5p [Saccharomyces cerevisiae]	4.6
		(U53340) coded for by C. elegans cDNA yk39e8.5; weakly similar to	
·····	255865	C. elegans proteins F09G8.4 and F02E8.6	3.6
1337 2	687999	(AE001123) B. burgdorferi predicted coding region BB0110	3.3
		(L38896) This CDS feature is included to show the translation of the	T PATRICULAR STATE
***************************************		corresponding V_region. Presently translation qualifiers on V_region	*
1338 6		features are illegal	3.1
1339 9	2000 AAAAAAAAAAAAAAAAAAAA	(X65335) lacZ [Cloning vector pSV-beta-Galactosidase Control]	2.7
1340 1	136390	(D79986) similar to human DNA-binding protein 5. [Homo sapiens]	2.6
1341 3	599395	(AF083072) histone H1 DNA binding protein [Cenarchaeum	1.4
		(U34781) Antho-LWamidII preprohormone [Anthopleura	·
1342 1			1.2
1343 3			0.98
1344 3	132825	(AF063403) putative cytosine-5 DNA methyltransferase [Zea mays]	0.89
1345 1	754989	(U47661) proline-rich protein PRP2 precursor [Lupinus luteus]	0.078
1346 7		!!!! ALU SUBFAMILY SB2 WARNING ENTRY	0.075
	ernamonare e e encommentario encommentario encomentario encomentario encomentario encomentario encomentario en	(U42436) C49H3.3 gene product [Caenorhabditis elegans]	0.023
	at the contract of the contrac	(AF053768) brain specific cortactin-binding protein CBP90 [Rattus	1e-009
	****	sodium bicarbonate cotransporter 2 >gi 3097316 dbj BAA25898	ュレーリンプ
1349 4:	507029	(AB012130) sodium bicarbonate cotransporter [Homo sapiens]	50.016
1350 2	781381	(AC004013) Similar to rabbit A-kinase-anchoring protein sapiens	5e-016
1351 4:	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(AJ131693) AKAP450 protein [Homo sapiens]	1e-029
1359 13	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U66260) multidrug resistance related protein 1	3e-052
1362 34	***************************************	( ) TO CO ! ! C	5.2
1363 1	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	TT 10 10 () 0 10 TYO 0	6.8
1505 1	······································	U42436) C49H3.3 gene product [Caenorhabditis elegans]	0.001
1364 1	10760	COAGULATION FACTOR X PRECURSOR 3.4.21.6) precursor -	
1367 38	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		2
130/138	DOLLLI (	AJ235272) unknown [Rickettsia prowazekii]	5.5

SEQ ID	Nearest Neighb ACCESSION	or (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VALU
SEQ ID	ACCESSION	CYTOCHROME P450 1A1 (CYPIA1) rainbow trout >gi 213780	IF VALU
1270	117143	(M21310) cytochrome P450IA1	7.1
13/0	11/143	HYPOTHETICAL 36.3 KD PROTEIN IN NRDC-MOBD	7.1
1271	3915910	INTERGENIC REGION >gi 1667570 (U76612) unknown	5.4
······································	1708230	HOMEOBOX PROTEIN ABDOMINAL-A	4.1
	3024292	RHODOPSIN >gi 2290717 (AF000947) rhodopsin [Sepia officinalis]	
13/3	3024292		)
1274	2996206	(AF053723) region 2 capsular polysaccharide biosynthesis protein [Actinobacillus pleuropneumoniae]	2.3
and the second s	2228750	[Actinobacinus pieuropheumomae] (U93868) RNA polymerase III subunit [Homo sapiens]	5e-011
13/3	2220130	hypothetical 30.5K protein - Enterococcus faecalis plasmid pAM-beta	1
1270	79960	1 >gi 3023044 (AF007787) orfC	2e-025
13/7	79900	HYPOTHETICAL 14.2 KD PROTEIN IN BLAB 3'REGION	; ZC-UZJ
		>gi 282541 pir  C41855 orf3 - Streptomyces cacaoi	Water
1290	165522		. 7
13004	465532	>gi 217001 dbj BAA00776  (D00937) regulatory protein for beta- COAT PROTEIN PRECURSOR (CAPSID PROTEIN) >gi 535774	5.7
1201	2012270	` , , ,	0.00
1381	3913279	(L09205) capsid protein [Tobacco ringspot virus] (AL034368) predicted using hexExon; L779.1, Serine/threonine	0.66
1		protein kinase, len: 1359 aa; Similarity to protein kinases. A.thaliana	
1290	4402763	NPK1-related protein kinase (TR:O22041) BLAST score: 303, sum	1.6
	4493762 131002		4.6
	632098	PROLINE-RICH PROTEIN MP-3 >gi 200549	0.1
1391	032098 	isp3 protein - fission yeast HYPOTHETICAL 6.1 KD PROTEIN C03B1.10 IN	4.5
1202	2406062		C- 00C
	2496862	CHROMOSOME X >gi 1072242 (U40952) C03B1.10 gene product	6e-006
13931.	3851703	(AF100421) p80 [Rattus norvegicus] (AL034368) predicted using hexExon; L779.1, Serine/threonine	4e-056
andinates in		· · · · · · · · · · · · · · · · · · ·	f 
1200	4493762	protein kinase, len: 1359 aa; Similarity to protein kinases. A.thaliana NPK1-related protein kinase (TR:O22041) BLAST score: 303, sum	
	1 <b>296</b> 48	PAIRED BOX PROTEIN PAX-1	4.3 4.3
1400;	129040	(D89861) cytochrome C-type biogenesis protein CCMF	4.3
1401.	4115789	[Cyanidioschyzon merolae]	1.9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	135514	T-CELL RECEPTOR BETA CHAIN PRECURSOR precursor (ANA	
1402	133314	(U41264) coded for by C. elegans cDNA CEESN26F; coded for by	0.034
Management of the Control of the Con		C. elegans cDNA CEESI89F; similar to 60S acidic ribosomal protein	
1402	1086833	Po (L10) [Caenorhabditis elegans]	2- 000
	4104168		3e-009
140414	+104100	(AF033339) UNC-45 [Caenorhabditis briggsae] HYPOTHETICAL PROTEIN HI1452 Haemophilus influenzae	7e-013
1410	1175805	<del>-</del>	7.2
ana ang ang ang ang ang ang ang ang ang	1708230	**************************************	7.3 4.2
1411	1708230	MYOSIN II HEAVY CHAIN, NON MUSCLE heavy chain	4.2
1412	107774	· · · · · · · · · · · · · · · · · · ·	0.56
<del>andrews and the second and the seco</del>	127774 728833	[Dictyostelium discoideum] !!!! ALU SUBFAMILY SB1 WARNING ENTRY	0.56
was a succession of the succes	and the second s		0.48
1414	3080645	(AC004611) Hsp27 ERE-TATA-binding protein [Homo sapiens]	3e-008
1/15	3687476	(AL031786) putative atp dependent rna helicase [Schizosaccharomyces pombe]	10.014
1417	306/4/0		1e-014
i i		down-regulated in adenoma protein down-regulated in adenoma	
Witespengellini	100	(DDA) human Sci 201064 576 500 570 500 : 4: 4	
1416	1557525	(DRA) - human >gi 291964 576-580, 579-583; acidic transcr. activ.	F - 0.00
etremenum et Metromorrament et magnion	and white the contract of the	domain 620-640,; homeobox motif 653-676 [Homo sapiens]	5e-060
1421 2	2088648	domain 620-640,; homeobox motif 653-676 [Homo sapiens] (AF002109) hypothetical protein [Arabidopsis thaliana]	6.9
1421 2 1422 1	2088648 1707768	domain 620-640,; homeobox motif 653-676 [Homo sapiens] (AF002109) hypothetical protein [Arabidopsis thaliana] (Y08256) orf c01038 [Sulfolobus solfataricus]	6.9 5.3
1421 2 1422 1	2088648 1707768	domain 620-640,; homeobox motif 653-676 [Homo sapiens] (AF002109) hypothetical protein [Arabidopsis thaliana] (Y08256) orf c01038 [Sulfolobus solfataricus] (AB008674) mBlm [Mus musculus]	6.9
1421 2 1422 1	2088648 1707768	domain 620-640,; homeobox motif 653-676 [Homo sapiens] (AF002109) hypothetical protein [Arabidopsis thaliana] (Y08256) orf c01038 [Sulfolobus solfataricus] (AB008674) mBlm [Mus musculus] (Z69663) Weak similarity to Cotton isocitrate lyase	6.9 5.3
1421 2 1422 1	2088648 1707768 3370996	domain 620-640,; homeobox motif 653-676 [Homo sapiens] (AF002109) hypothetical protein [Arabidopsis thaliana] (Y08256) orf c01038 [Sulfolobus solfataricus] (AB008674) mBlm [Mus musculus]	6.9 5.3

SEQ ID	ACCESSION	oor (BlastX vs. Non-Redundant Proteins)	
SEQ ID	ACCESSION	DESCRIPTION1	P VALU
1406	02006	hypothetical protein 1 (CYb-COII intergenic region) -	, Agreem
	83996	Sauroleishmania tarentolae mitochondrion uncertain [Leishmania	1.8
************	1322210	(U26347) immunogobulin kappa, VJ region [Homo sapiens]	1.4
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3043810	(U88273) NADH dehydrogenase subunit 4 [Sceloporus aeneus]	0.46
1428	3080645	(AC004611) Hsp27 ERE-TATA-binding protein [Homo sapiens]	0.0007
	deter reason	proteasome (prosome, macropain) 26S subunit, non-ATPase, 13	
***************************************	4506223	>gi 3618343 dbj BAA33214	1e-066
	2944430	(AF050157) butyrophilin-like [Mus musculus]	8.9
1439	1652823	(D90908) hypothetical protein	3
		EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-	The former country decreases printer-decreases
		RICH GLYCOPROTEIN) >gi 82169 pir  S06733 hydroxyproline-ricl	a 🏻
		glycoprotein precursor - common tobacco >gi 19867 emb CAA32090	1
979/8609 MANAGEMENT (979/75/24)	119714	(X13885) extensin (AA 1-620) [Nicotiana tabacum]	0.9
1441	509813	(L29010) ORFB [Cryphonectria hypovirus]	0.78
1442	238617	(S64572) nonfibrillar collagen Partial, 907 aa] [Strongylocentrotus	0.2
1443	3522970	(U42390) Trio [Homo sapiens]	0.12
1449	3328726	(AE001303) ATP Synthase Subunit E [Chlamydia trachomatis]	8.6
1450	501174	(L33965) MHC class II protein [Morone saxatilis]	8.5
***************************************	2661685	(AL009199) hypothetical protein SC7B7.10	8.5
	1575684	(U70316) IonA [Dictyostelium discoideum]	erigen variation and the commence of the comme
		hypothetical protein MJ0749 - Methanococcus jannaschii	8.4
1453	2128472	>gi 1591462 (U67520) conserved hypothetical protein	1
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1549329	(U52845) class IV chitinase EP3-1/H5 [Daucus carota]	5
	3845167	(AE001390) hypothetical protein [Plasmodium falciparum]	4.9
1.55		PUTATIVE IMPORTIN ALPHA SUBUNIT for by C. elegans	2.9
NO AND THE REST	***************************************	cDNA vk117h8 5; coded for by C. classers aDNA vk2000 5	i .
	***************************************	cDNA yk117h8.5; coded for by C. elegans cDNA yk8f10.5; coded	-
1456	2833306	for by C. elegans cDNA yk134a4.5; coded for by C. elegans cDNA	**
1430	and the second s	yk53c4.5; coded for by C. elegans cDNA yk8f10.3; coded for by C.	1.7
1462		HYPOTHETICAL 34.2 KD PROTEIN F07F6.2 IN	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		CHROMOSOME III >gi 746449 (U23486) No definition line found	8.3
·		(AF063232) variant 1 major surface glycoprotein [Pneumocystis	8.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(AC007154) hypothetical protein	4.8
anne e consequence de la conseque de	139372	ENDOPROTEASE (LATE L3 23 KD PROTEIN) adenovirus type 7]	4.8
1467	mananananananananananananananananananan	(X04813) CAD protein (AA 1 - 2236) [Drosophila melanogaster]	3.6
************************************		myosin-binding protein H - chicken	2e-005
1472	265108	(S54379) sucrose-phosphate synthase, SPS oleracea]	2.1
		HYPOTHETICAL 73.3 KD PROTEIN C6G9.14 IN	
1473	2842704	CHROMOSOME I >gi 1644326 emb CAB03616.1  pr otein	1.6
Transportation of the Contraction of the Contractio		NEURONAL ACETYLCHOLINE RECEPTOR PROTEIN, NON-	
-	4	ALPHA-3 CHAIN PRECURSOR (GFN-ALPHA-3) non-alpha-3	
1474	13109	chain precursor - goldfish >gi 212956 auratus]	0.7
and the same of th	(	down-regulated in adenoma protein down-regulated in adenoma	
a and a	(	DRA) - human >gi 291964 576-580, 579-583; acidic transcr. activ.	
1475 4	557535	1	1e-051
1485 2	315339 (	A FO1 (420)	9.9
	Ī	HYPOTHETICAL 23.0 KD PROTEIN IN IXR1-TFA1	~ • /
PROBLEM		NTERGENIC REGION >gi 539152 pir  S37847 hypothetical protein	
area property.		YKL030w - yeast (Saccharomyces cerevisiae)	
1486 5	49664	-1/49C0201 1 ICH 1010C41 (700000)	4.4
encontraction and and acceptance	204234 (	AFO25270\ MADCI	4.4
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	ORICRIN >gi 110649 pir  A35628 loricrin - mouse >gi 198871	0.51
1488 1	26390	M2.4200\ 1'- '- '- 5\ 4	0.14
		1123-300 Total III [19105 Husculus] >g1 220480 (UU9189) Ioricrin	0.14

SEQ ID		oor (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VAL
***************************************		down-regulated in adenoma protein down-regulated in adenoma	II VAL
	- Marian	(DRA) - human >gi 291964 576-580, 579-583; acidic transcr. activ.	
1489	4557535	domain 620-640,; homeobox motif 653-676 [Homo sapiens]	4e-050
******************	1707768	(Y08256) orf c01038 [Sulfolobus solfataricus]	4.3
~~~~	4376203	(U35226) putative cytochrome P-450	
POPER PROPERTY AND ADDRESS OF THE PARTY AND ADDRESS.	3721884	(AB016248) sterol-C5-desaturase [Mus musculus]	4.3
***************************************		ACIDIC PROLINE-RICH PROTEIN PRECURSOR salivary protein	4.2
1495	131000	[Rattus norvegicus]	1
	3123638	(AJ223069) TCF-3 protein [Mus musculus]	3.3
MARKET CONTRACTOR OF THE PARTY	854065	(X83413) U88 [Human herpesvirus 6]	0.49
17/1	1007000	GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2	5e-010
	Managari Jan	ANTIGEN) (ICE DINDING PROTEIN) (25 KD LECTER)	
	97-	ANTIGEN) (IGE-BINDING PROTEIN) (35 KD LECTIN)	
1501	126670	(CARBOHYDRATE BINDING PROTEIN 35) (CBP 35)	
1301	126679	GALACTOSIDE-BINDING LECTIN) >gi 1072481 pir  A28651	0.21
1500	4106560	(Z83819) dJ146H21.2 (similar to CYTOCHROME B-245 HEAVY	***
***********************	4106562	CHAIN) [Homo sapiens]	1e-011
· · · · · · · · · · · · · · · · · · ·	265108	(S54379) sucrose-phosphate synthase, SPS oleracea]	1.6
1505	1177322	(X95466) CPG2 protein [Rattus norvegicus]	0.0002
		(Z99942) similar to von Willebrand factor type A domain; cDNA	
		EST yk412d4.5 comes from this gene; cDNA EST yk412d4.3 comes	
manamaron on second	3878057	from this gene [Caenorhabditis elegans]	0.039
······	2394509	(AF024503) No definition line found [Caenorhabditis elegans]	6.1
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2827872	(U79229) NADH dehydrogenase subunit F [Tradescantia zebrina]	4.6
<b>~~~~</b>	3004653	(AF017777) tweety [Drosophila melanogaster]	3.7
1512	3873773	(Z83316) Similarity to S.pombe hypothetical protein C2F7.02C	1.6
all Awaren		HYPOTHETICAL 33.8 KD PROTEIN IN YSW1-RIB7	
apr-sc den		INTERGENIC REGION >gi 626526 pir  S46023 hypothetical protein	
71.00 (M. C.)		YBR152w - yeast (Saccharomyces cerevisiae)	Yaman
	586295	>gi 536480 emb CAA85111  (Z36021) ORF YBR152w	0.081
1519	3882147	(AB018256) KIAA0713 protein [Homo sapiens]	9.9
	2117181	(Z95584) mcr [Mycobacterium tuberculosis]	7.3
1521	1169862	G-BOX BINDING FACTOR 3 >gi 600863 thaliana]	6
1522	3257950	(AP000006) 236aa long hypothetical protein	6
1523	4138677	(AJ009814) polymerase [Viral hemorrhagic septicemia virus]	5.9
1524	2736413	(AF039044) No definition line found [Caenorhabditis elegans]	5.7
1525	2058691	(U94836) ERPROT 213-21 [Homo sapiens]	2.7
1526	347124	(L20967) phosphodiesterase [Homo sapiens]	2
- Company		FERRIC REDUCTASE TRANSMEMBRANE COMPONENT 3	
1,660cccoppe		PRECURSOR >gi 2132959 pir  S67293 probable membrane protein	AND
V WOTERAND		YOR381w - yeast (Saccharomyces cerevisiae)	1
			2
1527	3913674	He deponds of the control of the con	2
1527		>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w	2
	**************************************	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide	**************************************
1528	1086636	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans]	0.9
1528	1086636 2495706	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo	**************************************
1528 1 1529 2	1086636 2495706	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein	0.9 0.86
1528 1529 1530	1086636 2495706	Sgi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein Schizosaccharomyces pombe]	0.9 0.86 0.65
1528 1 1529 2 1530 4 1531 1	1086636 2495706 4007775 113671	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein Schizosaccharomyces pombe] !!! ALU CLASS F WARNING ENTRY !!!!	0.9 0.86 0.65 0.39
1528 1 1529 2 1530 4 1531 1 1532 4	1086636 2495706 4007775 113671 4098124	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein Schizosaccharomyces pombe] !!! ALU CLASS F WARNING ENTRY !!!! U73522) AMSH [Homo sapiens]	0.9 0.86 0.65 0.39 8e-008
1528 1 1529 2 1530 4 1531 1 1532 4 1533 1	1086636 2495706 4007775 113671 4098124 1174546	Sgi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein (Schizosaccharomyces pombe] !!! ALU CLASS F WARNING ENTRY !!!! U73522) AMSH [Homo sapiens]	0.9 0.86 0.65 0.39 8e-008 5e-010
1528 1 1529 2 1530 4 1531 1 1532 4 1533 1 1534 4	1086636 2495706 4007775 113671 4098124 1174546 1139228	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein Schizosaccharomyces pombe] !!! ALU CLASS F WARNING ENTRY !!!! U73522) AMSH [Homo sapiens] SYNAPTOTAGMIN II AF109134) 7-60 [Homo sapiens] >gi 4139272 (AF112980) 7-60	0.9 0.86 0.65 0.39 8e-008 5e-010 5e-044
1528 1 1529 2 1530 4 1531 1 1532 4 1533 1 1534 4 1535 2	1086636 2495706 4007775 113671 4098124 1174546 1139228 (2564328	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein Schizosaccharomyces pombe] !!! ALU CLASS F WARNING ENTRY !!!! U73522) AMSH [Homo sapiens] SYNAPTOTAGMIN II AF109134) 7-60 [Homo sapiens] >gi 4139272 (AF112980) 7-60 AB006628) KIAA0290 [Homo sapiens]	0.9 0.86 0.65 0.39 8e-008 5e-010 5e-044 3e-050
1528   1529   2 1530   4 1531   1 1532   4 1533   1 1534   4 1535   2 1536   3	1086636 2495706 4007775 113671 4098124 1174546 1139228 (2564328 6668141	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein Schizosaccharomyces pombe] !!! ALU CLASS F WARNING ENTRY !!!! U73522) AMSH [Homo sapiens] SYNAPTOTAGMIN II AF109134) 7-60 [Homo sapiens] >gi 4139272 (AF112980) 7-60 AB006628) KIAA0290 [Homo sapiens] AJ007398) PBK1 protein [Homo sapiens]	0.9 0.86 0.65 0.39 8e-008 5e-010 5e-044 3e-050 3e-057
1528   1529   2 1530   4 1531   1 1532   4 1533   1 1534   4 1535   2 1536   3 1544   3	1086636 2495706 	>gi 1420821 emb CAA99713  (Z75289) ORF YOR381w (U41013) similar to phosphoribosylaminoimidazolecarboxamide formyltransferase [Caenorhabditis elegans] HYPOTHETICAL PROTEIN KIAA0136 product is novel. [Homo AL034486) hypothetical fungal binuclear cluster domain protein Schizosaccharomyces pombe] !!! ALU CLASS F WARNING ENTRY !!!! U73522) AMSH [Homo sapiens] SYNAPTOTAGMIN II AF109134) 7-60 [Homo sapiens] >gi 4139272 (AF112980) 7-60 AB006628) KIAA0290 [Homo sapiens] AJ007398) PBK1 protein [Homo sapiens]	0.9 0.86 0.65 0.39 8e-008 5e-010 5e-044 3e-050

ible Z	B Nearest Neigh	ibor (BlastX vs. Non-Redundant Proteins)	
more recommendation of the	ACCESSION		P VAL
POPEL MORE DE LO COMPANY DE LA	6 3983015	(AF096056) antigen receptor [Ginglymostoma cirratum]	3.2
*********	7 1552169	(D42138) PIG-B [Homo sapiens]	3
~~~~	8 2440180	(Z99531) ubiquitin system protein	2.4
154	9 3882159	(AB018262) KIAA0719 protein [Homo sapiens]	1.8
155	0 3004653	(AF017777) tweety [Drosophila melanogaster]	1.3
155	1 77657	hypothetical 30.1K protein - Pseudomonas aeruginosa	0.26
155	2 4098124	(U73522) AMSH [Homo sapiens]	0.002
		(Z49967) cDNA EST EMBL:T00743 comes from this gene; cDNA	0.002
		EST EMBL:D69356 comes from this gene; cDNA EST	
	1000	EMBL:D65790 comes from this gene; cDNA EST EMBL:D70463	1
155	3 3979818	comes from this gene; cDNA EST EMBL:D66620 comes from this	0.001
	4 3550295	(AJ009947) putative ATPase [Homo sapiens]	
<del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	5 4507507	UNKNOWN >gi 3929583 (AF098162) timeless homolog [Homo	1e-008
***********		(AC002986) Contains similarity to membrane associated salt-	1e-035
156	2 3152567	inducible protein obligations of the E0072 (0 from A 11 1)	
1302	2 3132307	inducible protein gb/AF007269 from A. thaliana.	8.4
1561	3 4402079	(AL034559) predicted using hexExon; MAL3P7.21 (PFC0960c),	
Compared a second and compared to the compared	3 4493978	Hypothetical protein, len: 1929 aa	5
1304	1169862	G-BOX BINDING FACTOR 3 >gi 600863 thaliana]	4.7
	Nederland of the Control of the Cont	HYPOTHETICAL 259 KD PROTEIN (ORF 2136)	1
		>gi 81341 pir  A05037 hypothetical protein 2136 - liverwort	
CONTRACTOR OF THE PROPERTY OF	140550	(Marchantia polymorpha) chloroplast polymorpha}	3.7
~~~~~~~~~~~	2224354	(AB001684) ORF104 [Chlorella vulgaris]	0.76
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	106322	hypothetical protein (L1H 3' region) - human	0.53
***************************************	3 3522937	(AC004411) unknown protein [Arabidopsis thaliana]	0.4
1569	4097433	(U61084) phorbolin 3 [Homo sapiens]	4e-017
1578	1944122	(AB002531) SSU1p [Saccharomyces cerevisiae]	9.7
1579	4127662	(Y11176) fructosidase [Cichorium intybus]	5.3
		EPENDYMIN I PRECURSOR (EPD-I) rainbow trout >gi 213412	
1580	119515	(M93697) ependymin	4.4
1581	1439625	(U64598) weakly similar to S. cervisiae PTM1 precursor	2.6
1582	2183261	(AF002133) MAV264 [Mycobacterium avium]	2.3
***************************************		(AC000098) ESTs gb AA042402,gb ATTS1380 come from this gene	
1583	2388564	[Arabidopsis thaliana]	1.9
1584	1381091	(U51723) V-SERA 1 [Plasmodium vivax]	9.8
		HYPOTHETICAL PROTEIN MG144 HOMOLOG Mycoplasma	J.0
1587	2496317	pneumoniae (SGC3) (ATCC 29342) >gi 1674380 genitalium	0.65
		HYPOTHETICAL PROTEIN MG144 HOMOLOG Mycoplasma	0.65
1588	2496317	pneumoniae (SGC3) (ATCC 29342) >gi 1674380 genitalium	0.64
*****	3184073	(AL023779) hypothetical protein	0.64
~~~~	3560165	(AL031525) hypothetical protein	8e-009
1070		RNA-DIRECTED RNA POLYMERASE (ORF1B) 2.7.7.48) - avian	3.7
			200
		infectious bronchitis virus (strain Beaudette) >gi 292953 (M95169)	***************************************
1612	122504	pol protein [Avian infectious bronchitis virus] >gi 331173 (M94356)	references
***************************************	133594	ORF 1b encodes a polypeptide of potential mol. wt. 300,000.	9.7
1013	3043622	(AB011121) KIAA0549 protein [Homo sapiens]	6.8
1614	72070	translation elongation factor eEF-1 alpha chain factor 1-alpha (AA 1 -	
······································	72870	461) [Mus musculus]	5.6
1015	2315365	(AF016441) No definition line found [Caenorhabditis elegans]	5.3
***************************************		TRICHOHYALIN >gi 539701 pir  A45973 trichohyalin - human	
	586120	>gi 292836 (L09190) trichohyalin	4.9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		nuclear antigen EBNA-3B - human herpesvirus 4	3.9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	281206	nacion unigen Ebita-3b - naman nerpesyllus 4	
1617	281206 2146731	FK506-binding protein - Arabidopsis thaliana	CONTRACTOR
1617	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	FK506-binding protein - Arabidopsis thaliana (Z81506) similar to Lectin C-type domain short and long forms, CUB	3.5

SEQ ID	ACCESSION	por (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	P VALU
		LACCASE 4 PRECURSOR precursor - Rhizoctonia solani	J P VALU
1620	2833191	>gi 1150568 emb CAA91042	12.2
***********************	2981221	(AF053091) eyelid [Drosophila melanogaster]	3.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1118179	(U42304) chitin synthase [Phytophthora capsici]	3.2
*************************	1351576	HYPOTHETICAL PROTEIN MG419 Mycoplasma genitalium	2.9
	320309	AL2 protein - beet curly top virus >gi 210683	2.9
	1523997	(X99510) CDK4/6 kinase [Drosophila melanogaster]	2.5
*************	3646450	(AL031603) conserved hypothetical protein.	2.1
CONTRACTOR OF THE PARTY OF THE	3599342	(AF081112) ORF2 [Mus musculus domesticus]	2
****************	2114473	(U96963) p140mDia [Mus musculus]	1.9
1020	21177 <i>J</i>	(AC004877) sco-spondin-mucin-like; similar to P98167 uncertain	1.5
1629	3638957	[Homo sapiens]	
	2558942		1
	3257709	(AF024714) interferon-inducible protein [Homo sapiens]	0.93
*************	346430	(AP000005) 115aa long hypothetical protein	0.69
~~~~~~~~~~~	4263788	adenylylcyclase type V-alpha - dog >gi 290082	0.46
1033	4203700	(AC006068) hypothetical protein	0.3
1624	2134199	myosin I beta - bullfrog >gi 602138 (U14549) myosin I beta [Rana	
1034	2134139	catesbeiana] >gi 603692 (U14382) myosin I beta [Rana catesbeiana]	0.044
1625	2134199	myosin I beta - bullfrog >gi 602138 (U14549) myosin I beta [Rana	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3776027	catesbeiana] >gi 603692 (U14382) myosin I beta [Rana catesbeiana]	0.036
1030	3770027	(AJ010475) RNA helicase [Arabidopsis thaliana]	2e-006
-		ATP-DEPENDENT RRNA HELICASE SPB4 cerevisiae]	
1637	134787	>gi 836754 dbj BAA09238  (D50617) rRNA helicase	
1037	134/0/	[Saccharomyces cerevisiae] SPB4 [Saccharomyces cerevisiae]	1e-015
season and		ATP-DEPENDENT RRNA HELICASE SPB4 cerevisiae]	
1629	134787	>gi 836754 dbj BAA09238  (D50617) rRNA helicase	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	4454698	[Saccharomyces cerevisiae] SPB4 [Saccharomyces cerevisiae]	4e-016
1037	7734070	(AF070661) HSPC005 [Homo sapiens] (AL033125) 1-evidence=predicted by content; 1-	2e-016
		method=genefinder;084; 1-evidence_end; 2-evidence=predicted by	ia .
1640	4185892	match; 2-match_accession=SWISS-PROT:P38205; 2-	
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	match_description=HYPOTHETICAL 77.9 KD PROTEIN IN	6e-021
1041.		(AB011084) KIAA0512 protein [Homo sapiens]	9e-041
16/13	2435574	(AF026209) similar to C. elegans olfactory receptor ODR-10	
version and the second	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(NID:g1235900) [Caenorhabditis elegans]	7.3
mannen mannen sen sen sen sen sen sen sen sen sen	······à	(Z82055) similar to Zinc finger, C4 type	5.7
1043		PUTATIVE FLAGELLA-RELATED PROTEIN C HYPOTHETICAL 64.3 KD PROTEIN IN RPS3 3'REGION	14.5
277			
		(ORF516) >gi 419731 pir  S34525 hypothetical protein 516 (rps3 3'	Manage of the second
1646	267478	region) - Euglena gracilis chloroplast >gi 2673852 emb CAA50104	
1647 8	настичностичностичности инсистементория дестичной и	(X70810) orf516; ttg start [Euglena gracilis]	2
1047		hypothetical 237 protein (psbA 5' region) - rye chloroplast (fragment)	0.49
1649 1	1255792	(U51993) similar drosophila lethal (2) giant larvae protien	
	~~~ <del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	(SP:Q08470) [Caenorhabditis elegans]	0.09
1003	·····	All-1 protein +GTE form - mouse (fragment)	9.6
1664 4		(D89861) cytochrome C-type biogenesis protein CCMF	
······································	······	Cyanidioschyzon merolae]	6.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		U31081) MnxB [Bacillus sp.]	6
	**************************************	The state of the s	5.8
			4.9
		nypothetical protein II (retroposon LINE-1)	4.8
1669 8	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	L10908) Gcap1 gene product [Mus musculus] marker protein [Mus	4.6
1670 1	160540	ARGININOSUCCINATE SYNTHASE argininosuccinate synthetase	
10/01	168540 (	argG) homolog - Haemophilus influenzae (strain Rd KW20)	4.5

	A CCEGGION	bor (BlastX vs. Non-Redundant Proteins)	
16/1	ACCESSION		P VALU
540900400400000000000000000000000000000	160385	(M63277) malaria antigen [Plasmodium falciparum]	<b>¦</b> 4
	1177607	(X92485) pva1 [Plasmodium vivax]	3.8
	2497678	ZYXIN >gi 1430883 emb CAA67510	2.5
***************************************	4049518	(AL031852) conserved hypothetical protein	2.1
1675	135514	T-CELL RECEPTOR BETA CHAIN PRECURSOR precursor (ANA	0.55
		LIPASE 1 PRECURSOR (TRIACYLGLYCEROL LIPASE)	M. P. C.
1676	729942	>gi 538817 pir  A47081 triacylglycerol lipase	0.45
***************************************		coagulation factor XII (Hageman factor)	
		>gi 119763 sp P00748 FA12_HUMAN COAGULATION FACTOR	Language
	order ressona	XII PRECURSOR (HAGEMAN FACTOR) (HAF) 3.4.21.38)	Anne v.
1677	4503629	precursor - human >gi 180357 (M17466) coagulation factor XII	0.27
~~~~~		CASPASE-5 PRECURSOR (ICH-3 PROTEASE) (TY PROTEASE)	ξ <b>υ.</b> Ζ/
		(ICE(REL)-III) enzyme ICErel-III - human >gi 903936 (U28015)	the state of the s
1678	1708392	cysteine protease [Homo sapiens]	
	3550295		4e-006
10/7	3330293	(AJ009947) putative ATPase [Homo sapiens]	1e-008
1600	2075400	(Z73906) cDNA EST EMBL:M88866 comes from this gene	
1080	3875400	[Caenorhabditis elegans]	1e-009
		(Z78539) Similarity to S.pombe hypothetical protein C4G8.04	
an and an		(SW:YAD4_SCHPO); cDNA EST EMBL:D27846 comes from this	***************************************
and the same of th		gene; cDNA EST EMBL:D27845 comes from this gene; cDNA EST	differences
1681	3874685	yk202h7.3 comes from this gene; cDNA EST yk202h7.5 come	6e-010
		(Z49967) cDNA EST EMBL:T00743 comes from this gene; cDNA	*
		EST EMBL:D69356 comes from this gene; cDNA EST	outone,
Michigan		EMBL:D65790 comes from this gene; cDNA EST EMBL:D70463	
1682	3979818	comes from this gene; cDNA EST EMBL:D66620 comes from this	2e-015
1683	4584877	(AF068302) choline/ethanolaminephosphotransferase [Homo	5e-020
		ASPARTATE CARBAMOYLTRANSFERASE 2 PRECURSOR	150-020
1685	3914503	(A CD A DELA EEL EEL AND	9.9
1686	1845893	(11(010)	9.7
***************************************	and the state of the second state of the second	HYPOTHETICAL 43.0 KD PROTEIN B0361.6 IN	, <b>9.</b> 1 
		CHROMOSOME III >gi 458954 (U00031) similar to H. marismortui	house, Vagge
1687	1730629	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	5e-014
	······································	gag polyprotein - red flour beetle retrotransposon Woot >gi 805076	136-014
1690	2133580	gas polyprotem and flour occur retrotratisposon woot >gr 0000/0	<del></del>
- 0,000		(LIOUSXA) (DRE)	
		(U09586) ORF 1	1.1
	126206	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	,
	126296	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang]	1.1
1695	126296	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat	6
1695 1696	126296 111816	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus]	5.3
1695 1696	126296 111816 1517923	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa]	6
1695 1696	126296 111816 1517923	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1-	5.3
1695 1696	126296 111816 1517923	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence end; 2-	5.3
1695 1696 1697	126296 111816 1517923	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match accession=AA264666; 2-	5.3
1695 1696 1697	126296 111816 1517923	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste	5.3 5.3
1695 1696 1697 1698 1699 1	126296 111816 1517923 2924552 1663706	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste	5.3 5.3 6e-012
1695 1696 1697 1698 1699 1	126296 111816 1517923 2924552 1663706 8879850	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder	6 5.3 5.3 6e-012 2e-014
1695 1696 1697 1698 1699 1	126296 111816 1517923 2924552 1663706 8879850	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder	5.3 5.3 6e-012
1695 1696 1697 1698 2 1699 1 1700 3	126296 111816 1517923 2924552 1663706 8879850	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder (AL023781) cytochrome c oxidase polypeptide v precursor	6 5.3 5.3 6e-012 2e-014 3e-034
1695 1696 1697 1698 1699 1700 3	126296 111816 1517923 2924552 1663706 3879850 3184085	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder (AL023781) cytochrome c oxidase polypeptide v precursor [Schizosaccharomyces pombe]	6 5.3 5.3 6e-012 2e-014 3e-034 8.7
1695 1696 1697 1698 1699 1 1700 3 1707 3 1708 3	126296 111816 1517923 2924552 1663706 3879850 3184085 3845206	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder (AL023781) cytochrome c oxidase polypeptide v precursor [Schizosaccharomyces pombe] (AE001400) novel protein kinase [Plasmodium falciparum]	6 5.3 5.3 6e-012 2e-014 3e-034 8.7 3.9
1695 1696 1697 1698 2 1699 1 1700 3 1707 3 1708 3 1709 2	126296 111816 1517923 2924552 1663706 3879850 3184085 8845206 283435	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder (AL023781) cytochrome c oxidase polypeptide v precursor [Schizosaccharomyces pombe] (AE001400) novel protein kinase [Plasmodium falciparum] hypothetical protein DGF-1 - Trypanosoma cruzi cruzi]	6 5.3 5.3 6e-012 2e-014 3e-034 8.7 3.9
1695 1696 1697 1698 2 1699 1 1700 3 1708 3 1709 2 1710 3	126296 111816 1517923 2924552 1663706 3879850 3184085 3845206 283435 3329623	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder (AL023781) cytochrome c oxidase polypeptide v precursor [Schizosaccharomyces pombe] (AE001400) novel protein kinase [Plasmodium falciparum] hypothetical protein DGF-1 - Trypanosoma cruzi cruzi] (AF078790) No definition line found [Caenorhabditis elegans]	6 5.3 5.3 6e-012 2e-014 3e-034 8.7 3.9 3 1.3
1695 1696 1697 1698 2 1699 1 1700 3 1708 3 1709 2 1710 3 1711 2	126296 111816 1517923 2924552 1663706 3879850 3184085 3845206 183435 3329623 1459999	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder (AL023781) cytochrome c oxidase polypeptide v precursor [Schizosaccharomyces pombe] (AE001400) novel protein kinase [Plasmodium falciparum] hypothetical protein DGF-1 - Trypanosoma cruzi cruzi] (AF078790) No definition line found [Caenorhabditis elegans] (AF013108) tubulin Uni3 [Chlamydomonas reinhardtii]	66 5.3 5.3 5.3 6e-012 2e-014 3e-034 8.7 3.9 3 1.3 0.32
1695 1696 1697 1698 2 1699 1 1700 3 1709 2 1710 3 1711 2 1712 1	126296 111816 1517923 2924552 1663706 3879850 3184085 3845206 283435 3229623 2459999 705447	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein [Nycticebus coucang] probable pol polyprotein-related protein 4 - rat >gi 56590 emb CAA37647  (X53581) ORF4 [Rattus norvegicus] (U51117) ascospore maturation 1 protein [Neurospora crassa] (AL022018) 1-evidence=predicted by content; 1- method=genefinder;084; 1-method_score=165.48; 1-evidence_end; 2- evidence=predicted by match; 2-match_accession=AA264666; 2- match_description=LD08351.5prime LD Drosophila melanogaste (D87685) similar to human transcription factor TFIIS (S34159). (Z81592) predicted using Genefinder (AL023781) cytochrome c oxidase polypeptide v precursor [Schizosaccharomyces pombe] (AE001400) novel protein kinase [Plasmodium falciparum] hypothetical protein DGF-1 - Trypanosoma cruzi cruzi] (AF078790) No definition line found [Caenorhabditis elegans] (AF013108) tubulin Uni3 [Chlamydomonas reinhardtii] BACTENECIN 7 PRECURSOR (BAC7) aries]	6 5.3 5.3 6e-012 2e-014 3e-034 8.7 3.9 3 1.3

<del>erom occurrence force (see payment</del>	ACCESSION		P VALUI
Section of the sectio	987050	(X65335) lacZ [Cloning vector pSV-beta-Galactosidase Control]	7.6
***********************	1334582	(X55026) ORF16; no ATG start codon	6.8
	2981631	(AB012223) ORF2 [Canis familiaris]	5.2
1727	628300	gene F protein - rinderpest virus virus]	4.8
		(Z77669) Similarity to Human aminopeptidase N	
		(SW:AMPN_HUMAN); cDNA EST EMBL:D36412 comes from this	
		gene; cDNA EST EMBL:D37688 comes from this gene; cDNA EST	3
1728	4008417	EMBL:D34550 comes from this gene; cDNA EST EMBL:D33568	3.9
		PROBABLE GLOBAL TRANSACTIVATOR transactivator-like	
1729	1170083	protein [Autographa californica nucleopolyhedrovirus]	3.4
		(AL022727) dJ80I19.7 (olfactory receptor-like protein (hs6M1-3))	
1730	3757727	[Homo sapiens]	3.3
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	······································
1731	126296	[Nycticebus coucang]	1.2
***************************************		LARGE STRUCTURAL PHOSPHOPROTEIN	
WWW.		PHOSPHOPROTEIN) (BPP) >gi 73955 pir  XPBEA9 large structural	en company
1732	130702	phosphoprotein pp150 - human cytomegalovirus phosphoprotein 150	0.97
***		EPITHELIAL CHLORIDE CHANNEL PROTEIN calcium-activated	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1706571	chloride channel [Bos taurus]	0.87
1734	2981631	(AB012223) ORF2 [Canis familiaris]	0.56
		(U55366) coded for by C. elegans cDNA yk85a2.5; coded for by C.	***************************************
1735	1280072	elegans cDNA yk85a2.3; Similar to cuticlin	0.44
		HYPOTHETICAL 6.1 KD PROTEIN C03B1.10 IN	
1736	2496862	CHROMOSOME X >gi 1072242 (U40952) C03B1.10 gene product	0.001
1737	2228750	(U93868) RNA polymerase III subunit [Homo sapiens]	1e-007
		UNKNOWN >gi 3264861 (U97670) eukaryotic translation initiation	***************************************
1738	4503511	factor eIF3, p35 subunit [Homo sapiens]	4e-031
*		UNKNOWN >gi 3264861 (U97670) eukaryotic translation initiation	
marina mariatetrana accessivações.	4503511	factor eIF3, p35 subunit [Homo sapiens]	1e-032
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	113667	!!!! ALU CLASS B WARNING ENTRY !!!!	5.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2736375	(AF039040) No definition line found [Caenorhabditis elegans]	6
1743	2736375	(AF039040) No definition line found [Caenorhabditis elegans]	3.4
WARREN AND AND AND AND AND AND AND AND AND AN		(U30261) G protein beta subunit-like; Method: conceptual translation	i
······································	1002672	supplied by author [Schistosoma mansoni]	5e-024
1752 4	4200151	(AJ011538) hypothetical protein virus]	3.9
		SLP1 PROTEIN (VACUOLAR PROTEIN SORTING PROTEIN 33)	·
		>gi 101624 pir  A34708 SLP1 protein SLP1 protein [Saccharomyces	
1753 1	134528	cerevisiae] >gi 173185	1.6
and the second of the second o		HYPOTHETICAL 37.7 KD PROTEIN C18B11.06 IN	
and the same of th	Words	CHROMOSOME I >gi 2130289 pir  S58305 hypothetical protein	
in and the second	1175386	CD L C10D 11 oc m	0.13
1756 2	2909528	(AL021932) PPE [Mycobacterium tuberculosis]	4.9
		(Z81531) cDNA EST EMBL:D66579 comes from this gene; cDNA	······································
- Automotive	MANAGEMENT	EST EMBL:D70408 comes from this gene; cDNA EST yk263d3.5	
	en e	comes from this gene; cDNA EST yk275c1.5 comes from this gene;	
1760 3	8876797	DALL FOR TAKEY GLOSES	3.9
1770 3		(A FOO((COA)) 1: 5 55	8.9
		CYTOKINE RECEPTOR COMMON BETA CHAIN PRECURSOR	
AAAAAAA 44000		>gi 110595 pir  A35782 cytokine receptor common beta chain	
1771 4			4.3
•••••	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
1772 1		Ta T	2.9
1773 7	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		2.9 1.9
	organis according recommendation of the contraction	ATTICITOR IN CONTRACTOR AND	0.3

SEQ ID	ACCESSION	DESCRIPTION1	P VALU
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
~~~~~~~~~~~~	126296	[Nycticebus coucang]	0.001
NAME AND ADDRESS OF THE PARTY O	3879850	(Z81592) predicted using Genefinder	2e-037
***************************************	4155993	(AE001560) putative [Helicobacter pylori J99]	1.4
\$78 / A A DOG \$1000 CONSTRUCTION OF THE PARTY OF THE PART	225047	reverse transcriptase related protein [Homo sapiens]	2.2
1790	3785977	(AC005560) putative growth regulator protein	9.4
	The second secon	(AP000005) 617aa long hypothetical prolyl endopeptidase	
	3257681	[Pyrococcus horikoshii]	7.8
1792	2317972	(U97553) capsid protein [murine herpesvirus 68]	1.5
		cytochrome-c oxidase (EC 1.9.3.1) chain I - Trypanoplasma borreli	
1793	1085738	mitochondrion (SGC6) >gi 563140 borreli]	0.014
		(AL034559) predicted using hexExon; MAL3P7.14 (PFC0925w),	***************************************
1794	4493973	Hypothetical protein, len: 489 aa	0.002
		CYTOCHROME C OXIDASE POLYPEPTIDE III honeybee	
		mitochondrion (SGC4) >gi 552443 (L06178) cytochrome oxidase	
1800	461796	subunit 3 [Apis mellifera ligustica]	9.1
1801	1850592	(U88295) carnitine palmitoyltransferase II [Rattus norvegicus]	9
1802	207854	(M35837) alpha-amylase/alpha-galactosidase fusion protein	8.7
		Ksp-cadherin - rabbit >gi 902886 (U28945) Ksp-cadherin	,
1803	2136989	[Oryctolagus cuniculus] cuniculus]	3.4
1804	288448	(X06487) bcl2-Ig fusion gene [Homo sapiens]	3.4
1805	1657752	(U62325) FE65-like protein [Homo sapiens]	2.1
·····	2981631	(AB012223) ORF2 [Canis familiaris]	5e-007
····	4502103	annexin 31 XXXI) [Homo sapiens]	1e-043
1808	2149830	(U92864) maturase [Quercus rubra]	9.6
	2981631	(AB012223) ORF2 [Canis familiaris]	0.046
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	728832	!!!! ALU SUBFAMILY SB WARNING ENTRY	0.035
	······································	(Z98547) predicted using hexExon; MAL3P3.17 (PFC0420w),	10.033
No.		Calcium-dependent protein kinase, len: 591 aa; Similarity to calcium-	1
de la constante de la constant		dependent protein kinases. P.falciparum calcium-dependent protein	10.00
1813	3649769	kinase CDPK2 (TR:O15865) BLAST Sco	2.1
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	Z.1
1814	126296		1.6
managan ranganada.	197968	(M26361) LINE/Ig H-chain fusion protein [Mus musculus]	1.6
1010		HYPOTHETICAL 6.1 KD PROTEIN C03B1.10 IN	0.036
1816	2496862	CHROMOSOME X >gi 1072242 (U40952) C03B1.10 gene product	0.001
·	135514	T-CELL RECEPTOR BETA CHAIN PRECURSOR precursor (ANA	0.001
·	·····	(AB011108) KIAA0536 protein [Homo sapiens]	(contract of the contract of t
	**************************************	!!!! ALU SUBFAMILY SX WARNING ENTRY	4e-016
······································	~~~~~	(X92485) pva1 [Plasmodium vivax]	7
marana marana marana di kacamatan di kacamatan di kacamatan di kacamatan di kacamatan di kacamatan di kacamata	Language of the commence of th	hypothetical protein (L1H 5' region) - human	0.23
erro-romania de completo de la completo del la completo de  la completo de la com		(AB012223) ORF2 [Canis familiaris]	0.071
*****************	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(U41272) Similar to man(9)-alpha-mannosidase.	2e-009
************************	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(AF007261) SecY-type transporter protein [Reclinomonas	1e-032
1027	*************************************	FMRFAMIDE-RELATED NEUROPEPTIDES PRECURSOR	2.9
1830 1		244 (200 (770 (40 )	0.007
1020	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	>gi 416208 (U03137) neuropeptide precursor FMRFamide-related	0.026
1834		LIPOATE-PROTEIN LIGASE A 6.3.4) A - Escherichia coli	
······································	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	/ · — - · · - · · · ·	7.1
·	announce consumer masses recommended	/	0.044
10414	·····		7.3
N/A/A/A		(AL034559) predicted using hexExon; MAL3P7.17 (PFC0940c),	
10/10		HIMOTOOTICAL protoin Ione VIII as	2
1842 4		(2	3 1.7

SEQ ID	ACCESSION	DESCRIPTION1	P VALU
		(AF015825) hypothetical ABC transporter [Bacillus subtilis]	***************************************
	Assertation	>gi 2633579 emb CAB13082  (Z99110) similar to amino acid ABC	5
1852	2612897	transporter (ATP-binding protein)	5.9
************	728836	!!!! ALU SUBFAMILY SP WARNING ENTRY	1.2
~~~~~~	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	0.17
************	728836	!!!! ALU SUBFAMILY SP WARNING ENTRY	0.05
***************************************	106323	hypothetical protein (L1H 5' region) - human	0.03
~~~~~~~~~	2981631	(AB012223) ORF2 [Canis familiaris]	~~å~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
1007		HEXOSE TRANSPORTER HXT8 YJL214w - yeast (Saccharomyce:	0.0002
		cerevisiae) cerevisiae] >gi 1015600 emb CAA89511  (Z49489) ORF	
1850	729785		
1037	129103	YJL214w [Saccharomyces cerevisiae]	0.12
1041	1701242	(U83119) ORF2 consensus sequence encoding endonuclease and	
Time commence and a party	1791243	reverse transcriptase minus RNaseH [Rattus norvegicus]	4e-006
1862	854065	(X83413) U88 [Human herpesvirus 6]	2e-006
10.55	150505	(U80931) strong similarity to class-III of pyridoxal-phoshate-	***
1863	1707274	dependent aminotransferases	3e-032
		(AF063866) ORF MSV145 hypothetical protein [Melanoplus	
1871	4049828	sanguinipes entomopoxvirus]	5.6
W. W		(U41538) weak similarity to the S. cerevisiae activator 1 05 KD	
1872	1109853	subunit (replication factor C 95 KD subunit)	4.5
1873	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	3.6
1874	1684995	(U20663) NADH dehydrogenase subunit [Encyclia tampensis]	1.6
1875	3599325	(AF081106) ORF1 [Mus musculus domesticus]	0.36
<del>,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,</del>	160379	(M63274) malaria antigen [Plasmodium falciparum]	0.16
······································	728837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	0.10
		myosin I beta - bullfrog >gi 602138 (U14549) myosin I beta [Rana	U.12
1878	2134199	catesbeiana] >gi 603692 (U14382) myosin I beta [Rana catesbeiana]	0.036
	1872200	(U22376) alternatively spliced product using exon 13A	0.036
		(U97012) contains similarity to a ground domain, also weakly similar	U.UU9
er coverage		to drosophila fork head domain transcription factor SLP1	· · · · · · · · · · · · · · · · · · ·
1880	1938524	(SP:P32030) [Caenorhabditis elegans]	0.17
amanana amanana amanana a	varanta de la composition della composition dell	(AF032382) metalloprotease-disintegrin [Xenopus laevis]	0.17
•••••	····	(AP032382) metalloprofease-disintegrin [Xenopus laevis] !!!! ALU SUBFAMILY SQ WARNING ENTRY	0.77
······································	***		0.33
·	728833	(X83413) U88 [Human herpesvirus 6]	1e-007
1000		!!!! ALU SUBFAMILY SB1 WARNING ENTRY	7.4
about the		PUTATIVE SERINE/THREONINE PROTEIN PHOSPHATASE	
1007	1252170	C27B7.6 IN CHROMOSOME IV serine/threonine protein	
		phosphatase [Caenorhabditis elegans]	7.2
and the second		!!!! ALU SUBFAMILY SQ WARNING ENTRY	7e-009
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	135942	(U04295) DNA-binding factor of bZIP class [Oryza sativa]	1
earrant are an annual agus			0.94
		(AF081114) ORF2 [Mus musculus domesticus]	0.55
manuscrape and the second		!!!! ALU SUBFAMILY SQ WARNING ENTRY	0.056
		(M22333) unknown protein [Homo sapiens]	0.002
		!!!! ALU SUBFAMILY SX WARNING ENTRY	0.004
	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	8
	3646450	(AL031603) conserved hypothetical protein.	7e-028
	2213560	(Z97052) hypothetical protein	5e-026
	3002527	4 h	0.066
····	072977	real control of the c	0.022
1907 7			0.019
1908 4		in the second	2e-005
The same of the sa	······································	(TIOO CEO)	2
		) F [	_

1915 1	4572297	(AF071799) T-cell surface glycoprotein CD4 precursor RIBOSOME RECYCLING FACTOR (RIBOSOME RELEASING	6
			ı
		FACTOR) (RRF) >gi 1361841 pir  A64248 ribosome releasing factor	r <del> </del>
	1350904	Mycoplasma genitalium (SGC3) releasing factor (frr) [Mycoplasma	3.7
1916 7	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	1.4
1917 7	728832	!!!! ALU SUBFAMILY SB WARNING ENTRY	0.0006
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	1
1918 1	126296	[Nycticebus coucang]	4e-005
		(U83119) ORF2 consensus sequence encoding endonuclease and	1
1919 1	791243	reverse transcriptase minus RNaseH [Rattus norvegicus]	4e-009
1920 1	CONTRACTOR	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	0.006
1921 3	· ·	ATPase 6 - Sauroleishmania tarentolae	7.7
		HYPOTHETICAL 61.8 KD PROTEIN C12B10.03 IN	·/·/
1923 1	723547	CHROMOSOME I >gi 1262416 emb CAA94693	0.6
	177607	(X92485) pva1 [Plasmodium vivax]	8.6
	068583	(AF000580) Rep-like [Dictyostelium discoideum]	4.9
	2120082	hypothetical protein II (retroposon LINE-1)	2.4
	335198		0.21
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	134328	(X03145) pot. ORF III [Homo sapiens]	0.0003
<del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	502377	ECH - chicken >gi 1037160 dbj BAA08364	8.3
<del></del>		B-cell growth factor 1 (12kD)	0.094
marriament and a second	417289	(U95740) Unknown gene product [Homo sapiens]	0.065
1933 5		(U14550) SThM [Homo sapiens]	0.006
1934 7		!!!! ALU SUBFAMILY J WARNING ENTRY	7e-006
***************************************	649741	(AJ000281) mucin [Homo sapiens]	5e-020
	551821	(AF058803) mucin 4 [Homo sapiens]	8e-021
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	519443	(AB017430) kinesin-like DNA binding protein [Homo sapiens]	3e-029
······································	710216	(U79260) unknown [Homo sapiens]	1e-005
1940 72	28837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	5e-008
100		(AL034559) predicted using hexExon; MAL3P7.32 (PFC1010w),	
1942 44		Hypothetical protein, len: 1322 aa	4.5
1943 22	20578	(D00570) open reading frame (251 AA) [Mus musculus]	0.077
prantingen of the second		(U28739) similar to alcohol dehydrogenase/ribitol dehydrogenase	1
1944 27	731377	[Caenorhabditis elegans]	1e-028
1945 45	587207	(AB020527) Na/PO4 cotransporter homolog	4e-034
1948 72	28832	!!!! ALU SUBFAMILY SB WARNING ENTRY	0.21
1949 92	29913	(V01442) ribosomal protein S8 [Xenopus laevis]	0.066
1950 72	28831	!!!! ALU SUBFAMILY J WARNING ENTRY	2e-006
	**************************************	(U42833) coded for by C. elegans cDNA cm16f6; coded for by C.	
-		elegans cDNA CEESU63F; similar to S. cerevisiae SOF1 protein	
1951 11	125754		1e-028
1952 45	***************************************	(AB020527) Na/PO4 cotransporter homolog	6e-047
1953 45	<del>๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛๛</del>	(AB020527) Na/PO4 cotransporter homolog	2e-051
1955 29		(AB012223) ORF2 [Canis familiaris]	0.027
1956 41	138064	(Y18301) reverse transcriptase [Ovine pulmonary adenocarcinoma	~*************************************
		zinc finger protein 140 (clone pHZ-39)	0.0002
		>gi 1731416 sp P52738 Z140_HUMAN ZINC FINGER PROTEIN	
		140 >gi 2136409 pir  C57785 zinc finger protein ZNF140 - human	
1961 45		***************************************	0.6
1701 43		>gi 487787 (U09368) zinc finger protein ZNF140 [Homo sapiens]	8.6
1062 17		SERINE/THREONINE PROTEIN KINASE SSK2 (SUPPRESSOR	
- 19D7E1/	11541	OF SENSOR KINASE 2) SSK2 gene product [Saccharomyces	5.9
		CALIBRADY AND CHILD I belo manadain [Anal-13	2
1963 32			3
	28836	!!!! ALU SUBFAMILY SP WARNING ENTRY	0.001 0.0002

CEO TO	A COTO	bor (BlastX vs. Non-Redundant Proteins)	
SEQ ID	** \$**********************************		P VALU
PPSN \$11001190010010500000000000000	2293566	(AF012896) ADP-ribosylation factor 1 [Oryza sativa]	2e-012
************************	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.091
1970	59977	(Z14310) tripartite fusion transcript PLA2L	2e-005
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	······································
	126296	[Nycticebus coucang]	9
1972	4155181	(AE001495) putative TYPE II RESTRICTION ENZYME	4.8
1973	2239204	(Z97209) vacuolar carboxypeptidase	0.008
1974	3342107	(AF075269) nef protein [Simian immunodeficiency virus]	6.9
***************************************		X-linked retinopathy protein (C-terminal, clone XEH.8c) - human	
1978	423149	(fragment) >gi 299471 bbs 129340 XEH.8c} [human, Peptide Partial	0.66
***************************************	***************************************	(U83119) ORF2 consensus sequence encoding endonuclease and	,
1979	1791243	reverse transcriptase minus RNaseH [Rattus norvegicus]	0.025
***************************************		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	(0.023
1980	126296	[Nycticebus coucang]	0.011
	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.011
	720031		0.0009
1082	1791243	(U83119) ORF2 consensus sequence encoding endonuclease and	
1702	1171243	reverse transcriptase minus RNaseH [Rattus norvegicus]	4e-006
1004	1903379	(U52073) differentially repressed by testosterone and	* table
1904	1903379	dihydrotestosterone [Mus musculus]	3.2
1005	1002270	(U52073) differentially repressed by testosterone and	
***************************************	1903379	dihydrotestosterone [Mus musculus]	3.1
	3599339	(AF081111) ORF2 [Mus musculus domesticus]	2e-007
1990	728835	!!!! ALU SUBFAMILY SC WARNING ENTRY	5.9
		CCAAT DISPLACEMENT PROTEIN (HOMEOBOX PROTEIN	
		CLOX) (CLOX-1) >gi 423173 pir  S33121 homeotic protein Clox -	White control to the
	729093	dog (fragment) >gi 2202 emb CAA48782	0.018
1992	126295	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	0.004
		(U42833) coded for by C. elegans cDNA cm16f6; coded for by C.	·
		elegans cDNA CEESU63F; similar to S. cerevisiae SOF1 protein	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1125754	(SP:P33750) [Caenorhabditis elegans]	1e-030
1994	3789771	(AF055386) MHC class I related protein precursor	1.9
1995	728832	!!!! ALU SUBFAMILY SB WARNING ENTRY	3.1
		FRUCTOSE-1,6-BISPHOSPHATASE fructose-1,6-bisphosphatase	<del>/</del>
1996	3023729	[Mus musculus]	0.21
		ADP-RIBOSYLATION FACTOR factor [Plasmodium falciparum]	
1997	3182916	>gi 1932731 (U57370) ADP-ribosylation factor [Plasmodium	2e-015
	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	1
man manananan kananga	728836	!!!! ALU SUBFAMILY SP WARNING ENTRY	0.0003
manage of the second	59977	(Z14310) tripartite fusion transcript PLA2L	7e-007
		laminin receptor homolog - mouse 295) [Mus musculus]	, / 6-00 /
2003	91035	>gi 228997 prf  1815216A laminin receptor [Cricetinae gen. sp.]	0.10
	, 1000	regulator of mitotic spindle assembly 1	0.12
2005	4506545		
***************************************	······································	>gi 1350799 sp P49646 RMS1_HUMAN REGULATOR OF	3.7
·····	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(U70935) reverse transcriptase [Peromyscus maniculatus]	0.015
HALLANDON CONTRACTOR AND		!!!! ALU SUBFAMILY SQ WARNING ENTRY	0.007
		!!!! ALU SUBFAMILY J WARNING ENTRY	0.0002
2009		!!!! ALU SUBFAMILY J WARNING ENTRY	5e-006
- Company		(AC002131) Similar to seryl-tRNA synthetase gb U10400 from S	
2010		cerevisiae. EST gb N96627 comes from this gene.	5e-029
		HYPOTHETICAL 6.1 KD PROTEIN C03B1.10 IN	************************************
	2496862	CHROMOSOME X >gi 1072242 (U40952) C03B1.10 gene product	0.17
2012	2443342	(D88764) alpha 2 type I collagen [Rana catesbeiana]	1.2

SEQ ID	ACCESSION	por (BlastX vs. Non-Redundant Proteins)  DESCRIPTION1	DATATI
~		(U64570) myelin/oligodendrocyte glycoprotein-16.3kD(a) precursor	P VAL
		[Homo sapiens] >gi 1546037 (U64571) myelin/oligodendrocyte	
2013	1546035	glycoprotein-16.3kD(b) precursor	10.15
<del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.15
······································	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	0.006
	728838		0.003
**************************************	728837	!!!! ALU SUBFAMILY SX WARNING ENTRY	0.0003
·····		!!!! ALU SUBFAMILY SQ WARNING ENTRY	1e-005
····	3002527	(AF010144) neuronal thread protein AD7c-NTP [Homo sapiens]	1e-005
en marine commercial consideration and the c	728838	!!!! ALU SUBFAMILY SX WARNING ENTRY	3e-005
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2879805	(AL021813) hypothetical protein	0.6
	1711659	TCP1-CHAPERONIN COFACTOR A taurus]	0.003
	113667	!!!! ALU CLASS B WARNING ENTRY !!!!	0.0007
2025	113667	!!!! ALU CLASS B WARNING ENTRY !!!!	0.0007
	,	hypothetical L1 protein (third intron of gene TS) - human	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	87765	>gi 364964 prf  1510254A L1 repetitive element ORF [Homo	0.0003
2027	4009460	(AF039401) calcium-dependent chloride channel-1 [Homo sapiens]	2e-011
		PUTATIVE CYSTEINYL-TRNA SYNTHETASE C29E6.06C	
~~~~~	1730840	(CYSTEINETRNA LIGASE) (CYSRS) YNL247w - yeast	2e-035
***************************************	4200165	(Y16262) neutral invertase [Daucus carota]	5.7
····	1150834	(U42471) Wiscott-Aldrich Syndrome protein homolog [Mus	0.032
2032	113669	!!!! ALU CLASS D WARNING ENTRY !!!!	0.002
2033	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.0008
2034	1657758	(U66707) densin-180 [Rattus norvegicus]	1e-024
2035	4165313	(AB022083) SOX30 protein [Homo sapiens]	0.42
2036	1657758	(U66707) densin-180 [Rattus norvegicus]	1e-024
2037	3193336	(AF069301) DBI-related protein [Homo sapiens]	7e-055
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	1
2038	126296	[Nycticebus coucang]	8e-008
2039 2	2981631	(AB012223) ORF2 [Canis familiaris]	6e-008
·····	1098124	(U73522) AMSH [Homo sapiens]	3e-018
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	9e-006
·····	728837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	5e-009
•••••••••••••••••••••••••••••••••••••••	······································	ZINC FINGER PROTEIN 142 (KIAA0236) Human zinc finger	126-009
2044	3123312	protein(ZNF142) [Homo sapiens]	0.7
2045 7		!!!! ALU SUBFAMILY J WARNING ENTRY	0.7
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		(U79260) unknown [Homo sapiens]	0.13
		(Z38113) incomplete orf, len: 160, CAI: 0.09 similar to MRP ECOL	0.005
2047 5			
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	~~~~~~~ <u>~</u>	P21590 39.9 KD PROTEIN	1e-035
		ORF [Simian virus 40]	1.6
2049 7	·····	!!!! ALU SUBFAMILY SX WARNING ENTRY	0.001
2050 7	CONTRACTOR OF THE PARTY AND ADDRESS OF THE PAR	!!!! ALU SUBFAMILY J WARNING ENTRY	3e-005
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	!!!! ALU SUBFAMILY J WARNING ENTRY	3e-005
		(AB012223) ORF2 [Canis familiaris]	6e-007
		(AB012223) ORF2 [Canis familiaris]	3e-007
enconomica de la companya de la comp		(U86758) netrin-2 like protein [Homo sapiens] sapiens]	5e-012
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		PROBABLE HISTONE DEACETYLASE 1-2	6e-031
*************************		60S RIBOSOMAL PROTEIN L5	0.015
2057 7		!!!! ALU SUBFAMILY SP WARNING ENTRY	0.0003
2058 1		!!!! ALU CLASS C WARNING ENTRY !!!!	0.001
	or o	(U64570) myelin/oligodendrocyte glycoprotein-16.3kD(a) precursor	**************************************
· · · · · · · · · · · · · · · · · · ·	****	[Homo sapiens] >gi 1546037 (U64571) myelin/oligodendrocyte	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	546035	glycoprotein-16.3kD(b) precursor	0.0006
2060 5			5.9
		!!! ALU SUBFAMILY SP WARNING ENTRY	~ - ~

		bor (BlastX vs. Non-Redundant Proteins)	
water concerns to his contract the sec	ACCESSION		P VALU
an anataranananan aran aran aran aran ar	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	6e-006
	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.013
<del>~~~~</del>	2072977	(U93574) putative p150 [Homo sapiens]	0.001
***************************************	2072972	(U93572) putative p150 [Homo sapiens]	8e-005
	130402	RETROVIRUS-RELATED POL POLYPROTEIN	4e-013
***************	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	0.002
*****************	728831	!!!!! ALU SUBFAMILY J WARNING ENTRY	0.007
***************************************	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	4e-006
2075	987050	(X65335) lacZ [Cloning vector pSV-beta-Galactosidase Control]	0.017
*****	106322	hypothetical protein (L1H 3' region) - human	7e-006
***************************************	804808	(M13100) unknown protein [Rattus norvegicus]	0.11
2079	347964	(L22453) TARBP-b gene product [Homo sapiens]	0.001
2080	1196432	[(M22333) unknown protein [Homo sapiens]	4e-005
2081	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	8e-006
2082	3023928	PROBABLE HISTONE DEACETYLASE 1-2	5e-037
		pleiotropic regulator 1 (PRL1, Arabidopsis homolog) >gi 2832296	30-037
2083	4505895	(AF044333) pleiotropic regulator 1 [Homo sapiens]	6e-071
acomic management and the second	728836	!!!! ALU SUBFAMILY SP WARNING ENTRY	2e-005
	······································	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	125-003
2087	126296	[Nycticebus coucang]	6e-008
***************************************	···········	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	06-009
2088	126296	[Nycticebus coucang]	7. 000
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	106322	hypothetical protein (L1H 3' region) - human	7e-006
	2981631	(AB012223) ORF2 [Canis familiaris]	2e-006
	3336903	(Y10809) bZIP DNA-binding protein	1e-014
	728831	!!!! ALU SUBFAMILY J WARNING ENTRY	2.1
-07-	, 2000 I	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	0.01
2093	126296	[Nycticebus coucang]	0.000
or and the second of the secon	728837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	0.002
mmeno-a-a-a-a-a-a-à-	113671	!!!! ALU CLASS F WARNING ENTRY !!!!	4e-008
	386786	(J04988) 90 kD heat shock protein [Homo sapiens]	0.0001
······	106322	hymothetical protein (1 111 2) main 1	3e-009
2071	100322	hypothetical protein (L1H 3' region) - human	7e-005
2000	126296	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	U5 100 100 100 100 100 100 100 100 100 10
	728831	[Nycticebus coucang]	0.1
	~~ <del>~~~~~~~~~~~</del>	!!!! ALU SUBFAMILY J WARNING ENTRY	1e-005
		(AB012223) ORF2 [Canis familiaris]	1e-006
2102	2300307	60S RIBOSOMAL PROTEIN L21 musculus]	4e-007
2102	4106562	(Z83819) dJ146H21.2 (similar to CYTOCHROME B-245 HEAVY	
	······································	CHAIN) [Homo sapiens]	2e-061
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		!!!! ALU SUBFAMILY J WARNING ENTRY	8e-006
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1869835	(Z86099) protein kinase [human herpesvirus 2]	8.9
· mariani mari	1754989	(U47661) proline-rich protein PRP2 precursor [Lupinus luteus]	6.7
· · · · · · · · · · · · · · · · · · ·	2580578	(AF000996) ubiquitous TPR motif, Y isoform [Homo sapiens]	0.0004
***************************************		40S RIBOSOMAL PROTEIN S15A	0.0003
		(AJ011081) mader-drop8 [Homo sapiens]	2e-006
		(AF066071) SP85; PsB [Dictyostelium discoideum]	1e-008
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	8e-009
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		(AF081110) ORF2 [Mus musculus domesticus]	5e-012
men men minimi men men algeria.	1185943	(Y17833) pol protein [Human endogenous retrovirus K]	2e-022
2118 5	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(X57960) ribosomal protein L7 [Mus musculus]	3e-027
	[]	HYPOTHETICAL 65.0 KD PROTEIN IN HUPB-COF	
2120 2	2506605	NTERGENIC REGION >gi 1580716 dbj BAA11649  hypothetical	4.5
	2642222 (	(AF030885) telomere-associated recQ-like helicase	

SEQ ID	ACCESSION	por (BlastX vs. Non-Redundant Proteins)	T =
oed in	ACCESSION	DESCRIPTION1	P VALU
2125	1085573	microtubule-associated protein MAP2 - rat molecular weight	
	3192899	microtubule-associated protein cell line, Peptide Partial, 381 aa]	1.9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	106322	[(AF066072) SP85; PsB [Dictyostelium discoideum]	0.024
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2981631	hypothetical protein (L1H 3' region) - human	1e-005
oma ma come contrata de la constitución de la const	403460	(AB012223) ORF2 [Canis familiaris]	2e-010
2129	403400	(L24521) transformation-related protein [Homo sapiens]	0.003
A. W. Carlotte Company		Nascent polypeptide associated complex alpha chain - human	
**************************************		>gi 556642 emb CAA56869  (X80909) Nascent polypeptide	
2130	1082633	associated complex alpha subunit [Homo sapiens] >gi 4092060	1
	1669472	(AF054187) alpha NAC [Homo sapiens]	3e-006
	1127036	(U53757) pol polyprotein [Feline immunodeficiency virus]	6.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	728835	(D26178) serine/threonine protein kinase	0.81
managaran ana aranga	728837	!!!! ALU SUBFAMILY SC WARNING ENTRY	2e-007
2130	720037	!!!! ALU SUBFAMILY SQ WARNING ENTRY	5e-008
2137	730246	OX-2 MEMBRANE GLYCOPROTEIN PRECURSOR precursor -	
	3341992	human (fragment) >gi 1335216 emb CAA28943	3e-008
www.woonourourourouries	1363325	(AF054174) histone macroH2A1.2 [Homo sapiens]	6e-033
·	3892705	RNA helicase HEL117 - rat >gi 897915	9.9
***************************************	2565196	(AL033545) putative protein [Arabidopsis thaliana]	0.14
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	106322	(AF000381) non-functional folate binding protein [Homo sapiens]	0.007
MANAGER OF SHARE SHOWING SHOW AND AND AND ADDRESS OF THE PARTY OF THE	106322	hypothetical protein (L1H 3' region) - human	5e-007
2173	100322	hypothetical protein (L1H 3' region) - human	0.033
2144	133902	40S RIBOSOMAL PROTEIN S27 ribosomal protein S27 isoform [Homo sapiens]	
***************************************	111814		0.0004
Market Market Colonia de la Sala	1669454	hypothetical protein 3 - rat norvegicus]	2e-011
2170	1009434	(U53748) pol polyprotein [Feline immunodeficiency virus] LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	8.6
2147 1	126296	[Nycticebus coucang]	
217/1	120270	ribosomal protein S23 S23 >gi 543449 pir  S41955 ribosomal protein	6e-005
- Address - Addr		S23 - rat protein [Homo sapiens] >gi 453281 emb CAA54584	Mentione
2148 4	1506701	(X77398) ribosomal protein S23 [Rattus norvegicus]	12 - 000
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	3e-009
2149 1		[Nycticebus coucang]	2- 011
interes and a contract of the	Parameter and a superior and a super	(U34998) Rad9 [Coprinus cinereus]	3e-011
in the second contract of the second contract	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	hepatocyte growth factor receptor - African clawed frog	4.8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	······································	(M22333) unknown protein [Homo sapiens]	3e-006
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	60S RIBOSOMAL PROTEIN L13A (23 KD HIGHLY BASIC	,36-000
		PROTEIN) >gi 345897 pir  S29539 basic protein, 23K - human	and an and an
2155 7		>gi 23691 emb CAA40254  (X56932) 23 kD highly basic protein	2e-008
management and the second second second	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(U93569) putative p150 [Homo sapiens]	3e-009
***************************************		(U83119) ORF2 consensus sequence encoding endonuclease and	36-009
2157 1	791243	reverse transcriptase minus RNaseH [Rattus norvegicus]	4e-012
		retinoblastoma-binding protein 1; RBP1	40-012
- Contraction of the Contraction		>gi 1710030 sp P29374 RBB1_HUMAN RETINOBLASTOMA	
at a constant	uspense	BINDING PROTEIN 1 (RBBP-1) >gi 2136103 pir  I58383	
2158 4		retinoblastoma binding protein 1 - human protein 1, RBP1 [human,	5e-008
·····		(AB012223) ORF2 [Canis familiaris]	0.0008
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	v.vvv0
2161 1		- · · · · · · · · · · · · · · · · · · ·	0.0005
2162 7	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	MEMBRANE-ASSOCIATED PROTEIN HEM-2 H19 protein -	2e-007
2163 1		(6)	20.012
	rancomence con conservation and a second and	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	2e-012
2165 12		37 1	60 005
		1	6e-005

SEQ ID	ACCESSION		P VALU
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
~~~~	126296	[Nycticebus coucang]	1e-012
2172 1	106323	hypothetical protein (L1H 5' region) - human	9e-010
99, 000		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
2173 1	26296	[Nycticebus coucang]	2e-013
	teritoria (de la composition de la comp	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	20-015
2174 1	26296	[Nycticebus coucang]	2e-013
2175 6	Maria managapapapapapapapapapapapapapapapapapap	(L39061) transcription factor SL1 [Homo sapiens]	4e-014
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	072964	(U93569) putative p150 [Homo sapiens]	~~~~ <del>``</del>
******************************	***************************************	microtubule-associated protein MAP2 - rat molecular weight	5e-007
2180 1	085573	microtubule-associated protein cell line, Peptide Partial, 381 aa	2.0
2181 1		collagen, corneal - chicken (fragment)	3.8
commence and the commence of t	981631	(AB012223) ORF2 [Canis familiaris]	0.097
·····	493820	CYTOCHROME C OXIDASE POLYPEPTIDE I ornatipinnis	1e-010
2103 2	773020	HYPOTHETICAL 71.7 KD PROTEIN F52H3.2 IN	3e-011
2184 2	494737		
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	709997	CHROMOSOME II >gi 3877390 emb CAA91322	3e-046
2186 5	MONANCE PROPERTY AND ALBERTA A	DNA REPAIR PROTEIN RAD18 pombe]	0.64
2180 3	***************************************	(Z14310) tripartite fusion transcript PLA2L	1e-006
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	**************************************	(U93570) putative p150 [Homo sapiens]	5e-007
2188 1		(U83883) p105 coactivator [Rattus norvegicus]	4e-010
2189 7		!!!! ALU SUBFAMILY J WARNING ENTRY	4e-009
2190 22	245560	(AF004339) cytochrome c oxidase subunit II [Homo sapiens]	9e-007
2101	055650	(Z77655) Weak similarity to Human calcium-dependent proetase	
2191 38		(SW:CANS_HUMAN)	0.2
2192 72	recommendation of the commence of	!!!! ALU SUBFAMILY SC WARNING ENTRY	2e-010
2193 10	**************************************	hypothetical protein (L1H 3' region) - human	3e-013
2194 20	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(U93570) putative p150 [Homo sapiens]	1e-015
2196 18	~~~~~~~ <i></i>	(M18728) ORF1 [Homo sapiens]	0.0005
2197 26	CONTRACTOR AND ADDRESS OF THE CONTRACTOR AND ADDRESS OF THE CONTRACTOR AND ADDRESS OF THE CONTRACTOR ADDRESS OF THE CONTRA	(AF027404) signal recognition particle 14a [Macaca radiata]	0.0003
2202 33	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(M80344) ORF2 contains a reverse transcriptase domain.	5e-017
2203 45	**************************************	(AL035526) extensin-like protein	1
2204 30	·	(AB011137) KIAA0565 protein [Homo sapiens]	3e-020
- Communication		HYPOTHETICAL 47.8 KD PROTEIN F57B9.5 IN	
2206 39	······································	CHROMOSOME III >gi 532824 (U13876) F57B9.5 gene product	0.1
2207 17	769472	(U15780) p82 [Homo sapiens]	9e-037
2208 72	28837	!!!! ALU SUBFAMILY SQ WARNING ENTRY	8e-015
2209 25	06089	26S PROTEASE REGULATORY SUBUNIT 7	1e-016
	orden van	aldehyde dehydrogenase (EC 1.2.1) aldB - Escherichia coli	***************************************
***************************************	discontinue de la contraction	>gi 912476 (U00039) No definition line found [Escherichia coli]	***************************************
2210 10	73466	>gi 1790014 (AE000436) aldehyde dehydrogenase B (lactaldehyde	1.4
2211 20	72960	(U93568) p40 [Homo sapiens]	3e-005
2212 40	09460	(AF039401) calcium-dependent chloride channel-1 [Homo sapiens]	2e-020
2214 10	0687	nydroxyproline-rich glycoprotein - rice glycoprotein [Oryza sativa]	8.7
2215 30	43616	AB011118) KIAA0546 protein [Homo sapiens]	2e-010
	······································	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	20 010
2216 12		Nycticebus coucang]	6e-014
2218 91	~~~~~	LINE-1 hypothetical protein - mouse (fragment) musculus]	3e-005
	***************************************	Z83819) dJ146H21.2 (similar to CYTOCHROME B-245 HEAVY	(2C-003
2220 410		CHAIN) [Homo sapiens]	50.061
2221 63		inc-containing protein - human	5e-061
2223 20	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U93572) putative p150 [Homo sapiens]	0.0001
2224 312	and the second s	HYPOTHETICAL PROTEIN	3e-009
T J 1	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		1e-012
2226 126		JNE-1 REVERSE TRANSCRIPTASE HOMOLOG protein Nycticebus coucang]	
	U42U	INYCHOODES COUCANY	1e-007

SEQ ID	Contract of the Contract of th		P VAL
222	7 2981631	(AB012223) ORF2 [Canis familiaris]	3e-013
	U MA L'ATANA	GTP-binding protein rah - mouse (fragment) G-protein=low-	***************************************
***************************************	8 109930	molecular-weight GTP-binding protein GTP-binding protein [Mus	4e-020
******************	9 2981631	[(AB012223) ORF2 [Canis familiaris]	5e-007
	1698455	(U49974) mariner transposase [Homo sapiens]	5e-007
223	1 2289235	(U95016) myocyte nuclear factor-beta [Mus musculus]	8e-015
		(AL009196) 1-evidence=predicted by content; 1-	100015
		method=genefinder;084; 1-evidence_end; 2-evidence=predicted by	40
		match; 2-match_accession=AA391048; 2-	AMERICAN AND AND AND AND AND AND AND AND AND A
	2827495	match_description=LD09991.5prime LD Drosophila melanogaster	1.9
2233	2465330	(U92818) unnamed HERV-H protein [Homo sapiens]	1e-011
2234	2135837	nuclear protein H731 - human >gi 1825562	4e-014
2235	4164448	(AF044958) NADH:ubiquinone oxidoreductase ASHI subunit	2e-025
**************************************		protein kinase C substrate 80K-H	120-023
		>gi 120629 sp P14314 G19P_HUMAN PROTEIN KINASE C	***
	The state of the s	SUBSTRATE, 80 KD PROTEIN, HEAVY CHAIN (PKCSH) (80K-	
2236	4506077	H PROTEIN) >gi 105167 pir  A32469 80K protein H precursor 80K-	4- 021
M94M200000000000000000000000000000000000	2981631	(AB012223) ORF2 [Canis familiaris]	
******	1176422	(U43194) rhophilin [Mus musculus]	0.007
***************************************		(AC004923) similar to UNC-93; similar to U89424 (PID:g3642687)	0.81
2239	4263743	[Homo sapiens]	2 014
CONTRACTOR OF STREET STREET STREET	106322	hypothetical protein (L1H 3' region) - human	3e-041
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	2352427	(AF004161) peroxisomal Ca-dependent solute carrier	2e-018
	4104400	(AF035401) exocellobiohydrolase precursor [Piromyces rhizinflata]	1e-044
~~~~	158154	(M81959) POU domain protein [Drosophila melanogaster]	4.2
		MAGNESIUM-PROTOPORPHYRIN IX MONOMETHYL ESTER	3
		OVIDATIVE CVCLASE 66 KD SUDLAHTED 1.1.1	
2244	114858	OXIDATIVE CYCLASE 66 KD SUBUNIT Rhodobacter capsulatus >gi 46113 emb CAA77530  (Z11165) 575 aa (66 kD) oxidative	
***************************************	2352427	(AF004161) peroxisomal Ca-dependent solute carrier	2.2
		PROBABLE COAT PROTEIN virus >gi 58812 emb CAA29526	3e-044
2246	116746	(X06166) ORF IV (AA 1-489)	
	2072967	(U93570) putative p150 [Homo sapiens]	4.3
		M-sema F protein precusor - mouse homolog=M-Sema F [mice,	6e-008
2248	2137494		*
	1709971	60S RIBOSOMAL PROTEIN L10A (CSA-19)	5.5
·····	1498225	(V07560) DILADIO	1e-015
2230	1470223	(Y07569) PHAPI2a protein [Homo sapiens] TRANSLOCON-ASSOCIATED PROTEIN, DELTA SUBUNIT	4e-012
7		PRECURSOR (TRAD DELTA) (SIGNAL GEOLIENGE PROTEGOR	
li diamento de la constanta de	www.	PRECURSOR (TRAP-DELTA) (SIGNAL SEQUENCE RECEPTOR	
2251	1711550	DELTA SUBUNIT) (SSR-DELTA) unnamed protein product [Homo	
2231	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	sapiens] >gi 1302656 sapiens] >gi 1673433 emb CAA92215	3e-012
2252	126296	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
2232	······································	[Nycticebus coucang]	1e-013
Contraction		chaperonin containing T-complex subunit 6	
2252	1502642	>gi 730922 sp P40227 TCPZ_HUMAN T-COMPLEX PROTEIN 1,	
······································	4502643 2737894	ZETA SUBUNIT (TCP-1-ZETA) (CCT-ZETA) (TCP20) sapiens]	1e-014
	····	(U59151) Cbf5p homolog [Homo sapiens]	4e-070
	1076557	extensin-like protein - cowpea (fragment)	0.6
MMMMM	annapproxima .	(Z69664) Similarity to Yeast cell divison control protein cdc25	
**************************************	e et enem	(SW:CC@%_SACKL); cDNA EST EMBL:D32475 comes from this	
2267	1070245	gene; cDNA EST EMBL:D34376 comes from this gene; cDNA EST	
2257 3	8878245	EMBL:D35124 comes from this gene; cDNA EST EMBL:D37	9.2
MM Parket	1  505067	MAD2 (mitotic arrest deficient, yeast, homolog)-like 1 >gi 950199 U31278) mitotic feedback control protein Madp2 homolog [Homo	······································
		1101000	

EQ ID	ACCESSION	DESCRIPTION1	P VAL
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
2259	126296	[Nycticebus coucang]	4e-018
		tumor protein, translationally-controlled 1 CONTROLLED TUMOR	
		PROTEIN (TCTP) (P23) factor - human >gi 37496 emb CAA34200	
2260	4507669	(X16064) tumor protein (AA 1 - 172) [Homo sapiens]	0.043
and the state of t		arylamine N-acetyltransferase (EC 2.3.1.5), monomorphic - rabbit	·
		>gi 217728 dbj BAA00989  (D10108) arylamine N-acetyltransferase	;
2261	66499	[Oryctolagus cuniculus]	1.8
***************************************	733532	(U23420) unknown [Drosophila melanogaster]	0.51
***************************************	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	DEOXYURIDINE 5'-TRIPHOSPHATE	0.31
		NUCLEOTIDOHYDROLASE PRECURSOR (DUTPASE) (DUTP	
		PYROPHOSPHATASE) >gi 2443580 (AF018432) dUTPase [Homo	***************************************
2263	3041664	sapiens] >gi 2735292 (U90223) deoxyuridine triphosphate	20 020
management and a second	2136246	tastin - human >gi 905356 (U04810) tastin	2e-029
******	126295	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	1.1
······································	825671		0.0002
····	2231019	(X16934) B23 nucleophosmin (280 AA) [Homo sapiens]	4e-013
market market market market market	-	(Z97207) B-IND1 protein [Mus musculus]	2e-023
2209	4502189	aquaporin 8 sapiens]	1e-014
2270	12/20/	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
2270	126296	[Nycticebus coucang]	5e-015
2271	10/00/	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
	126296	[Nycticebus coucang]	5e-015
TOTAL PROPERTY OF THE PROPERTY AND THE PROPERTY AND	218564	(D90358) HB-SOD [Schizosaccharomyces pombe]	4e-024
~~~~	1352736	PELOTA PROTEIN >gi 973224 (U27197) pelota [Drosophila	3e-038
~~~~~~~~	102177	hypothetical protein (ribosomal RNA repeat region) - Giardia lamblia	PONT NEW WORLDSON, MAN PROPERTY AND
2276	113668	!!!! ALU CLASS C WARNING ENTRY !!!!	0.13
		(D63876) KIAA0154 gene product is related to mouse gamma	
manamanamanaman	961444	adaptin. [Homo sapiens]	7e-026
annon annon anno anno	3183217	HYPOTHETICAL PROTEIN KIAA0103 sapiens]	5e-017
2282	2072972	(U93572) putative p150 [Homo sapiens]	4e-017
C. Market		(D63876) KIAA0154 gene product is related to mouse gamma	
····	961444	adaptin. [Homo sapiens]	6e-028
2285	4586287	(AB004794) DUF140 [Xenopus laevis]	0.005
Alleganija		(Z74042) predicted using Genefinder; Similarity to Haemophilus 3-	
ONTON Asses		oxoacyl-(acyl-carrier protein) reductase (SW:FABG_HAEIN); cDNA	and the same of th
uuu.		EST yk470b2.3 comes from this gene; cDNA EST yk470b2.5 comes	
2286	3879684	from this gene [Caenorhabditis elegans]	0.0002
		RAS-RELATED PROTEIN RAB-1A protein ypt1 - mouse	
in the same of the		>gi 2144599 pir  TVHUYP GTP-binding protein Rab1 - human	x seems
-		>gi 2144600 pir  TVDGYP GTP-binding protein Rab1 - dog	
		>gi 55457 emb CAA68284  (Y00094) Ypt1 protein (AA 1-205) [Mus	
2287	131786	musculus] >gi 550060 (M28209) GTP	1e-017
2288	4115532	(AB012043) NBR13 [Homo sapiens]	0.69
2289	4586287	(AB004794) DUF140 [Xenopus laevis]	1e-007
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	
2290	126296	[Nycticebus coucang]	9e-020
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	······	Pathogenesis-Related Protein 5d From Nicotiana Tabacum	7.8
*******************************	······································	cell surface antigen CD34 precursor - human	1e-019
			0.14
vannamatara a a a a a a a	interpretation that we arrest an enterpretation accommends to the	(L30113) alcohol dehydrogenase; ADH [Papio hamadryas]	2e-018
******	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	(U57715) FGF receptor activating protein FRAG1 [Rattus	1e-020
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	collagen, type IV, alpha 4 CHAIN PRECURSOR	020
		>gi 1360674 pir  CGHU1B collagen alpha 4(IV) chain precursor -	
	ĝ.	ZEILTOOOLAIDII ICCUTO LD COHASCH SIOHS #H L L LUSIU DISCUISOF = 3	

		or (BlastX vs. Non-Redundant Proteins)	I Day a za z
SEQ ID	Constitution of the contract o	DESCRIPTION1	P VALUI
2299	2565196	(AF000381) non-functional folate binding protein [Homo sapiens]	1e-006
		NEURONAL ACETYLCHOLINE RECEPTOR PROTEIN, ALPHA-	,
		6 CHAIN PRECURSOR >gi 1458118 (U62435) nicotinic	4 4
		acetylcholine receptor alpha6 subunit precursor [Homo sapiens]	
2300	2492620	>gi 2815225 emb CAA76155  precursor [Homo sapiens]	1e-009
		hypothetical protein 2 - Agrobacterium tumefaciens	
~~~~~~	95095	>gi 39102 emb CAA37890.1  (X53945) ORF2	3.3
	1765938	(U47077) DNA-dependent protein kinase catalytic subunit	6e-021
A STOCK AND THE WORLD BY A STOCK AND THE	2493783	COLLAGEN ALPHA 4(IV) CHAIN bovine (fragment)	3
THE PROPERTY OF THE PROPERTY OF THE	2645205	(U63648) p160 myb-binding protein [Mus musculus]	3e-029
**********	2645205	(U63648) p160 myb-binding protein [Mus musculus]	9e-030
2306	2495322	HOMEOBOX PROTEIN HOX-A9	8e-014
		CALCYCLIN (PROLACTIN RECEPTOR ASSOCIATED	Open son
		PROTEIN) (PRA) (GROWTH FACTOR-INDUCIBLE PROTEIN	
2307	116509	2A9) (S100 CALCIUM-BINDING PROTEIN A6) put. calcyclin;	4e-006
		cytochrome c oxidase subunit VIIb	
		>gi 461804 sp P24311 COXM_HUMAN CYTOCHROME C	
		OXIDASE POLYPEPTIDE VIIB PRECURSOR 1.9.3.1) chain VIIb	-
2308	4502991	human >gi 30151 emb CAA78613  sapiens]	0.002
		tumor protein, translationally-controlled 1 CONTROLLED TUMOR	
		PROTEIN (TCTP) (P23) factor - human >gi 37496 emb CAA34200	
2309	4507669	(X16064) tumor protein (AA 1 - 172) [Homo sapiens]	6e-020
2311	4507207	sorcin sorcin [Homo sapiens] >gi 1094394 prf  2106141A sorcin	2e-018
		CYTOCHROME C OXIDASE POLYPEPTIDE III chain III - human	
2312	117061	mitochondrion (SGC1) oxidase III [Homo sapiens]	3e-017
2313	1709972	60S RIBOSOMAL PROTEIN L10A (CSA-19)	5e-013
2314	4454698	(AF070661) HSPC005 [Homo sapiens]	3e-014
2315	1885381	(U77665) RNaseP protein P30 [Homo sapiens]	1e-020
2316	209383	(M27786) MS-2 pol-stefin B fusion protein [Artificial gene]	7e-008
2318	631507	zinc-containing protein - human	0.03
		cytochrome c oxidase subunit VIIb	
		>gi 461804 sp P24311 COXM_HUMAN CYTOCHROME C	
		OXIDASE POLYPEPTIDE VIIB PRECURSOR 1.9.3.1) chain VIIb -	
2319	4502991	human >gi 30151 emb CAA78613  sapiens}	5e-013
		hypothetical L1 protein (third intron of gene TS) - human	
2320	87765	>gi 364964 prf  1510254A L1 repetitive element ORF [Homo	2e-015
		MITOCHONDRIAL MATRIX PROTEIN P1 PRECURSOR (P60	
Î		LYMPHOCYTE PROTEIN) (60 KD CHAPERONIN) PROTEIN)	
e de la companya de l		(HUCHA60) >gi 107086 pir  A32800 heat shock protein 60 precursor	
2321	129379	- human >gi 190127 (M22382) mitochondrial matrix protein [Homo	8e-016
2322	2231019	(Z97207) B-IND1 protein [Mus musculus]	6e-022
2323	4154176	(U96639) ATPase subunit 6 [Canis familiaris]	0.007
2324	2575807	(D49692) adenylate cyclase [Spirulina platensis]	4.4
		cytochrome c oxidase subunit VIIb	\$1000000000000000000000000000000000000
		>gi 461804 sp P24311 COXM_HUMAN CYTOCHROME C	
, dominate and a second		OXIDASE POLYPEPTIDE VIIB PRECURSOR 1.9.3.1) chain VIIb	
2325	4502991	·	5e-007
	***************************************	P43 - human >gi 833999 bbs 160014 (S75463) P43=mitochondrial	***************************************
2326	2119918		3e-020
	ka magamanaga managa ann 60 agam harap majankaga pama nga mbakaga pinda asa pada k	cathepsin E precursor - human >gi 181194 (J05036) cathepsin E	and the second second
2327	4503145	precursor [Homo sapiens] >gi 181205 (M84424) cathepsin E	1e-041
and the second s	4502189		2e-023
~~~~~	4502189	aquaporin 8 sapiens]	2e-026
entrantemente de conservações	2306969	(AF007860) xl-Mago [Xenopus laevis]	2e-037
4330	<b>ムン</b> ひひプ <b>じ</b> ブ	(VI AN 100A) VI-IATARA [VEITOBRE 196A12]	2C-U3/

EQ ID	ACCESSION	DESCRIPTION1	P VAL
<b>2</b> 333	3970731	(AJ006480) SBT4B protein [Lycopersicon esculentum]	9
		(AF039052) Similar to inosine monophosphate dehydrogenase;	***************************************
		coded for by C. elegans cDNA CEMSF04F; coded for by C. elegans	
		cDNA yk247b12.3; coded for by C. elegans cDNA cm20d8; coded	5 5 8
2334	2736524	for by C. elegans cDNA yk247b12.5; coded for by C. elegans	2.7
44-04-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0	~ ····	hypothetical protein Y - Streptomyces nogalater	
2335	1085957	>gi 2147591 pir  S69232 hypothetical protein Y	1.1
********************	3183217	HYPOTHETICAL PROTEIN KIAA0103 sapiens]	3e-021
		TRANSLATION INITIATION FACTOR EIF-2B GAMMA	30-021
2337	2494312	SUBUNIT (EIF-2B GDP-GTP EXCHANGE FACTOR) subunit	3e-037
***************************************	337930	(M22146) scar protein [Homo sapiens]	1e-024
	68891	transforming protein ets - chicken >gi 211753	2e-007
		LINE-1 REVERSE TRANSCRIPTASE HOMOLOG protein	26-007
2340	126296	[Nycticebus coucang]	0- 017
25.0	120270	tight junction protein 1 (zona occludens 1)	9e-017
		>gi 585098 sp Q07157 ZO1_HUMAN TIGHT JUNCTION	
		PROTEIN ZO-1 (TIGHT JUNCTION PROTEIN 1) human	
23/1	4507517		
<del>~~~</del>	106851	>gi 292938 (L14837) tight junction (zonula occludens) protein ZO-1	9e-023
2343	100631	keratin 18, cytoskeletal - human (fragment) sapiens]	4e-023
2244	1702420	HYPOTHETICAL 52.3 KD PROTEIN C56F8.06C IN	*
eroromentor con incompe	1723438	CHROMOSOME I PRECURSOR	7.5
2343	2231019	(Z97207) B-IND1 protein [Mus musculus]	7e-022
2246	2047200	(L40459) latent transforming growth factor-beta binding protein [Mus	1
2346	2047300	musculus]	0.48
Management		PROBABLE RNA-DEPENDENT HELICASE P68 human	
***************************************		>gi 35220 emb CAA36324  (X52104) p68 protein (AA 1-614) [Homo	account of the contract of the
		sapiens] >gi 38318 emb CAA33751  (X15729) protein p68 (AA 1-	
2347	129383	614) [Homo sapiens] >gi 2599360	2e-023
		BCL2-interacting killer BCL-2 INTERACTING KILLER	1
·····	4502411	(APOPTOSIS INDUCER NBK) (BP4) sapiens] >gi 1235989	3e-025
****	2072961	(U93568) putative p150 [Homo sapiens]	2e-014
	2306969	(AF007860) xl-Mago [Xenopus laevis]	3e-041
2355	1872498	(U74297) PiUS [Oryctolagus cuniculus]	2e-033
***************************************		(AB000170) endopeptidase 24.16 type M3 endopeptidase 24.16 type	1
podes (mesong		M3 [Sus scrofa] type M3 [Sus scrofa] >gi 1783130 dbj BAA19065	100000
arramana arramana da a	1783123		1e-029
entero anacemano ele-	2306969	(AF007860) xl-Mago [Xenopus laevis]	2e-041
	1872498	(U74297) PiUS [Oryctolagus cuniculus]	6e-034
	4519602	(AB017563) IGSF4 [Homo sapiens]	3e-041
mmmmmmmmmmän.	1944330	(D49545) KIFC2 [Mus musculus]	2e-019
2361	4502189	aquaporin 8 sapiens]	6e-040
		aquaporin 8 sapiens]	2e-041
	1537070	(U63840) nucleoporin p54 [Rattus norvegicus]	2e-040
2364	3024124	HOMEOBOX PROTEIN MEIS3	8e-030
2365			5e-040
2366 2		(AF007860) xl-Mago [Xenopus laevis]	7e-054
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	/	2e-054
	······	(AC007136) Putative map kinase interacting kinase [Homo sapiens]	
			7e-045
2368 4		I DOMO Sabiensi	
2368 4	1464284	[Homo sapiens] cyclin-dependent kinase 6 KINASE 6 (KINASE PLSTIRE)	/C-U+J
2368 4	1464284	cyclin-dependent kinase 6 KINASE 6 (KINASE PLSTIRE)	/C-U+J
2368 4	1464284 i		/ C-U+3

		or (BlastX vs. Non-Redundant Proteins)	
SEQ ID	ACCESSION	DESCRIPTION1	P VALU
		HYPOTHETICAL 68.1 KD PROTEIN B0304.7 IN	
2371	2496815	CHROMOSOME II >gi 1041881 (U39472) similar to f44f4.5	0.59
	-	HYPOTHETICAL 68.1 KD PROTEIN B0304.7 IN	
2372	2496815	CHROMOSOME II >gi 1041881 (U39472) similar to f44f4.5	0.56
2373	3327226	(AB014606) KIAA0706 protein [Homo sapiens]	1e-031
*****************************	3327226	(AB014606) KIAA0706 protein [Homo sapiens]	2e-034
	3327226	(AB014606) KIAA0706 protein [Homo sapiens]	2e-038
2376	2231019	(Z97207) B-IND1 protein [Mus musculus]	2e-039
2377	3327226	(AB014606) KIAA0706 protein [Homo sapiens]	4e-043
		NEUROLYSIN PRECURSOR (NEUROTENSIN	***************************************
		ENDOPEPTIDASE) (MITOCHONDRIAL OLIGOPEPTIDASE M)	
		ANGIOTENSIN-BINDING PROTEIN) (SABP) protein - pig	
		>gi 217709 dbj BAA01949  (D11336) soluble angiotesin-binding	
2378	417743	protein [Sus scrofa] type M1 [Sus scrofa] >gi 1871389 dbj BAA1	1e-047
	***************************************	(AB000172) endopeptidase 24.16 type M2 endopeptidase 24.16 type	1
2379	1783127	M2 [Sus scrofa]	7e-050
······································		TRANSLATION INITIATION FACTOR EIF-2B GAMMA	
2380	2494312	SUBUNIT (EIF-2B GDP-GTP EXCHANGE FACTOR) subunit	7e-056
	1085499	CDC4 repeat unit-containing protein - mouse	4e-056
2382	4092846	(AB019987) chromosome-associated polypeptide-C [Homo sapiens]	4e-058
		(AF023265) NAD+-specific isocitrate dehydrogenase beta subunit	1
2383	4103446	isoform A [Homo sapiens]	5e-059
2384	1353709	(U42385) FIN16 gene product [Mus musculus]	8e-061
	······	(AF083395) phospholipase A2-activating protein [Homo sapiens]	5e-061
2386	473407	(U08215) NST-1 [Mus musculus]	2e-061
		(AF023265) NAD+-specific isocitrate dehydrogenase beta subunit	
2387		isoform A [Homo sapiens]	6e-062
2388	1537070	(U63840) nucleoporin p54 [Rattus norvegicus]	2e-062
2389	3851584	(AF092563) chromosome-associated protein-E [Homo sapiens]	7e-063
2390		The state of the s	6e-063
2391	3493209		3e-064
		cyclin-dependent kinase 6 KINASE 6 (KINASE PLSTIRE)	······································
Distriction.		>gi 107662 pir  S23387 protein kinase (EC 2.7.1.37) cdk6 - human	
WWW. arve		>gi 4389286 pdb 1BI8 A Chain A, Mechanism Of G1 Cyclin	
2392		Dependent Kinase Inhibition From The Structures Cdk6-P19ink4d	5e-065
2393		(AF052577) aldo-keto reductase [Homo sapiens]	7e-066
2394		/	3e-068
2395		(AF083395) phospholipase A2-activating protein [Homo sapiens]	1e-068
2396			1e-070

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALU
101	U73106	Liriodendron tulipifera high-pI laccase (LAC2-4) mRNA,	4.9
and the same of th		Bacillus thuringiensis insecticidal crystal protein (CryIF)	4
manne	M63897	gene, complete cds.	4.8
103	X75014	M.musculus Phox2 mRNA for homeodomain protein	4.4
ACCRECATE AND ADDRESS OF A STATE OF	X77320	A.officinalis L. unknown mRNA	4.1
annemara necessariam na és	M64982	Human fibrinogen alpha chain gene, complete mRNAs.	<sup>1</sup> 4.1
	U28241 ·	Gallus gallus collapsin-3 mRNA, partial cds.	4.1
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M36860	Human elastin mRNA, complete cds.	4.1
~~~~	X99641	M.musculus mRNA for HP1 alpha protein	4.1
109	X92557	S.erythraea pccB, bcpA2, and orfX genes	4
110	M21514	Rinderpest virus fusion protein mRNA, complete cds.	4
111	M64982	Human fibrinogen alpha chain gene, complete mRNAs.	4
112	D87218	Trypanosoma cruzi mRNA, partial cds, clone:TcEST002	4
	***************************************	E.coli rafA, rafB, and rafD genes encoding alpha-D-	
113	M27273	galactosidase, raf-permease, and raf-invertase, complete cds.	3.9
114	U50065	Caenorhabditis elegans cosmid T23A7.	3.9
115	X76186	Rinderpest virus (RBOK vaccine) mRNA for matrix protein	3.9
116	L05165	Gorilla gorilla glycophorin-gene related sequence.	3.9
***************************************		Apis koschevnikovi mitochondrial cytochrome oxidase II	\$~~~~~~~ !
117	M77212		3.9
		Human ELK1 pseudogene (ELK2) and immunoglobulin	***************************************
118	U64453	heavy chain gamma pseudogene (IGHGP)	3.9
**********	illione mantee de conservation de la conservation d	Rotavirus SA114F nonstructural protein 34 (NS34) mRNA,	\$0000000000000000000000000000000000000
119	M87502	complete cds.	3.9
		Bacillus subtilis ribosomal protein L7/12 (rplL) gene, beta	<u> </u>
120	L24376	· · · · · · · · · · · · · · · · · · ·	3.9
	U97143	опростичения по принципального по принцения выправания выправания выправания выправания выправания выправания в	3.9
***	M95610		3.9
······································	M21890	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3.9
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Homo sapiens integrin, alpha 5 (fibronectin receptor, alpha	<del>.</del>
		polypeptide) (ITGA5) mRNA > :: emb X06256.1 HSFNRA	) 6
124	NM 002205.1	Human mRNA for integrin alpha 5 subunit	3.9
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U38949	Gallus gallus cardiac C-protein mRNA, complete cds.	3.9
		Simian immunodeficiency virus clone SIVsm62J vpx gene,	ļ <del> </del>
126	U04985	partial cds, env, nef, tat and rev genes, complete cds.	3.9
	U05237		,3.9
	X96616	P.primaurelia gene encoding 156D surface antigen	3.8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U31929		3.8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X13523	Yeast CBS2 gene for cytochrome b translational activator	3.8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M57769	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3.8
	***************************************	HIV-1 isolate 652 clone 11 from Haiti, envelope	<b>3.</b> 0
132	U08443	glycoprotein (env) gene, partial cds.	3.8
	COOTTJ	Caldicellulosiruptor saccharolyticus alpha-dextrin 6-	J.U
133	L39876		3.8
anna ann an ann an ann an ann an an an a	M20363	and the contract of the contra	3.8
	J02836	aisanna Tanananan an a	3.8
Stateta cominences cominences	U64880	Mouse beta-glucuronidase gene, complete cds  Eubacterium thermomarinus ribonuclease P RNA	3.8
		······································	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X93520	E.caballus microsatellite DNA marker (clone ASB6)	3.8
138	Z83151		3.8
100	D20404	Aspergillus oryzae pgkA gene for phosphoglycerate kinase,	2.0
correction in the contract of the contract of the	D28484	<u> </u>	3.8
mananananananananananananananananananan	Z28091	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3.8
141	V01291	Yeast gene for alcohol dehydrogenase	3.8

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
	U44843	Lactococcus lactis plasmid pND324, complete sequence	3.8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M13496	Chicken type X collagen gene.	3.8
144	Y14946	Homo sapiens mRNA for SPIN protein	3.8
		Human muscle glycogen phosphorylase (PYGM) gene,	***************************************
	U94776	exons 6 through 17	3.8
	U43844	Mus musculus cyclin D3 gene, complete cds	3.8
	X03431	Drosophila melanogaster transposable element 297	3.8
148	U05237	Human fetal Alz-50-reactive clone 1 (FAC1) mRNA,	3.8
		Comptonia peregrina maturase (matK) gene, chloroplast	3
	U92856	gene encoding chloroplast protein, complete cds	3.8
150	X94165	Human papillomavirus type 73 E6, E7, E1, E2, E4, L2, and	3.7
		Drosophila azteca NDSSC 14012-0171.6 glycerolphosphate	
151	U47875	dehydrogenase (Gpd) gene, partial cds	3.7
152	X02882	Human HLA class II alpha chain gene DZ-alpha	3.7
153	AF005932	Clavispora opuntiae Spt3 (SPT3) gene, complete cds	3.7
154	Z11840	D.melanogaster hedgehog gene DNA	3.7
	700.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.000 00.00	Arabidopsis thaliana ecotype Landsberg K+ transport system	30000000 00000000000000000000000000000
155	U06745	AKT1 gene, complete cds.	3.7
	······································	Unidentified crenarchaeote 16S ribosomal RNA gene, 5'	
156	U63362	partial sequence	3.7
****		Wheat gene for transcription factor HBP-1b(c38), final	
157	D30810	exon, partial cds	3.7
*************	X56089	X. laevis mRNA for alpha-subunit of G-protein, type G-	3.7
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X07701	Chironomus tentans Balbiani ring mRNA BR 2.1 3'-end	3.7
M/Knaperawanan-outsianan-outsianan-	X64649	G.gallus mRNA for restrictin	3.7
<del></del>	Y13426	Homo sapiens TCRDV2 gene, partial	3.7
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Y14443		3.7
	U92794	Homo sapiens mRNA for zinc finger protein Mus musculus alpha glucosidase II beta subunit mRNA,	§
***************************************	Y09480	entermonarement mentermonarement and a comparement of the comparement	3.7
104	109460	A.europaeus genes encoding dehydrogenase and cytochrome	
		Homo sapiens ADP-ribosylation factor 3 (ARF3) mRNA > ::	
165	NIM 001650 1	gb M74491 HUMADPRF3A Human ADP-ribosylation	2.7
	NM_001659.1 L20893	factor 3 mRNA, complete cds.	3.7
······································		Rice yellow mottle virus complete genome.	3.7
16/	AF019759	Canis familiaris beta-glucuronidase (GUSB) mRNA,	3.7
1.00	1160505	Cricetulus griseus beta-1,6-N-acetylglucosaminyltransferase	
168	U62587		3.7
4.50		Cucumis sativus mRNA for NADPH-protochlorophyllide	
**************************************	D50085	oxidoreductase, complete cds	3.7
annon annon annon annon annon an Air	M81890	Human interleukin 11 (IL11) gene, complete mRNA.	3.7
171	M57765	Human interleukin 11 mRNA, complete cds.	3.7
		T. reesei ura3 (OMPdecase) gene for orotidine-5'-phosphate	
172	X55880	decarboxylase (EC 4.1.1.23)	3.6
3		P.falciparum dihydrofolate reductase-thymidylate synthase	
173	J03028	gene, complete cds.	3.6
174	AF000949	Canis familiaris keratin (KRT9) gene, complete cds	3.6
175	U78718	Dugesia tigrina 26S ribosomal RNA gene, partial sequence	3.6
176	D16471	Human mRNA, Xq terminal portion	3.6
177	X69838	H.sapiens mRNA for G9a	3.6
more movement receives accountly.	M24685		3.6
***************************************	L05468	Trichomonas vaginalis beta-tubulin (btub1) gene, complete	3.6
**************************************	Y08926	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3.6
	M59743	<i></i>	3.6
	X51952	**************************************	3.6

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
		Candida albicans phosphoglycerate kinase (PGK1) gene,	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U25180	complete cds.	3.6
184	X58080	Maize chloroplast ORF170 and psaA gene	3.6
		H. sapiens (D1S502) DNA segment containing (CA) repeat;	
185	Z24609	clone AFM361td9; single read	3.6
		Mus musculus (BALB/c) L1 repeat insertion in the kappa	ANY WARRACK
186	M24936	chain 5' flank.	3.6
		Thermoanaerobacterium thermosulfurigenes orfA gene,	Solomore
		partial cds, polygalacturonase precursor (pglA), abcA, abcB	***
187	U50951	and sigma factor (sigA) genes, complete cds	3.6
188	U57999	Mus musculus prosaposin (psap/SGP-1) gene, complete cds.	3.6
189	AF000949	Canis familiaris keratin (KRT9) gene, complete cds	3.6
**************************************		nucleoprotein [tomato chlorotic spot virus, isolate BR-03,	**************************************
190	S54325	Genomic RNA, 929 nt]	3.6
***************************************	***************************************	{endogenous retrovirus SY-3, provirus} [human,	•
191	S70572	lymphocytes, Genomic, 2189 nt]	3.6
***************************************	en e	Rhizobium sp. NGR234 plasmid pNGR234a, section 29 of	<del> </del>
192	AE000092	46 of the complete plasmid sequence	3.6
***************************************	U75285	Homo sapiens apoptosis inhibitor survivin gene, complete	3.6
	X91404	W.mirabilis mRNA for phosphoenolpyruvate carboxylase	3.6
	M17376	Mouse alpha-1-acid glycoprotein I (AGP-1) gene, complete	3.6
		Human DNA sequence from cosmid 92M18, BRCA2 gene	
196	Z73360	region chromosome 13q12-13	3.6
~~~~~	L05364		3.6
	J04353	Human papillomavirus type 31 (HPV-31) complete genome.	3.6
····	X03882	Paramecium primaurelia gene for G surface protein	3.6
~~~~	U53152	Caenorhabditis elegans cosmid K11D5.	3.6
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M65126	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	3.6
201	14103120	Yeast (S.cerevisiae) lysyl-tRNA synthetase (KRS1) alpha-2	2.0
202	J04186	subunit gene, complete cds.	3.6
202	30-100	Mus musculus (clone MKT6) morphogenetic protein 1	3.0
203	L35281	(BMP-1/His), alternative splice	3.6
	D83390	Gallus gallus mRNA for connectin/titin, partial cds	3.6
204	D03330	Glycine max partial SIRE-1 sequence gag-protease	3.0
205	U22103		2.6
203	U22103	polyprotein mRNA, complete cds Human interferon alpha2 upstream scaffold associated	3.6
206	U82705		2.6
200	082703	region 3 (SAR3) and non-SAR region DNA	3.6
202	T 120200	Rhagoletis cerasi 16S ribosomal RNA gene, mitochondrial	2.5
	U39389	4. Tarantana and an anti-anti-anti-anti-anti-anti-anti-anti-	3.5
	X13287	Medicago sativa nodulin-25 gene	3.5
	X91337	H.sapiens La/SS-B pseudogene 2	3.5
	M19684	Human alpha-1-antitrypsin-related protein gene, exons 3, 4	3.5
	M35296	Human tyrosine kinase arg gene mRNA.	3.5
manninen mannen manninen der Se	X88000	S.tenacellus mRNA for ubiquinol:cytochrome c	3.5
	M61906	Human P13-kinase associated p85 mRNA sequence.	3.5
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z29084	C.butyricum transposon containing tbcC gene	3.5
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M92039	decementation and the commence of the commence	3.5
216	D86478		3.5
Announce		Saccharomyces cerevisiae nuclear polyadenylated RNA-	
217	U35737	binding protein (NAB4) gene, complete cds.	3.5
TETTOWNWA.		B.thuringiensis 20 and 67 kd mosquitocidal protein genes,	
218	M22860	complete cds and IS231-like transposase, 3' end.	3.5

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALU
	and the same of th	H.sapiens CpG island DNA genomic Mse1 fragment, clone	
219	Z57857	1d10, forward read cpg1d10.f1a	3.5
	3900000000	Pneumocystis carinii clone PC14103 major surface	
	U07059	glycoprotein mRNA, complete cds.	3.5
	X52978	H.sapiens gene for lipoprotein lipase	3.5
222	M24566	Dictyostelium discoideum tRNA-Glu-GAA gene, clone	3.5
	0	Homo sapiens calcium dependent potassium channel alpha	
	U11058	subunit (MaxiK) mRNA, complete cds	3.5
	M21514	Rinderpest virus fusion protein mRNA, complete cds.	3.5
~~~~~~~~~	X80454	HIV type 1 (CA7) env-gene	3.5
	X98695	Bacteriophage T4 modA and modB genes	3.5
COMMERCIA COMPANION CONTRACTOR AND	X76186	Rinderpest virus (RBOK vaccine) mRNA for matrix protein	3.5
	S82456	PGHS-2=prostaglandin endoperoxide H synthase-2	3.5
	D85194	Arabidopsis thaliana mRNA, partial cds	3.5
******************************	L35661	Homo sapiens (subclone H8 4_b9 from P1 35 H5 C8) DNA	3.5
231	L76205	Colletotrichum gloeosporioides non-LTR retrotransposon	3.5
		Homo sapiens xeroderma pigmentosum, complementation	Account Actor
		group F (XPF) mRNA > :: gb U64315 HSU64315 Human	
232	NM_000464.1	DNA repair endonuclease subunit	3.5
233	X02155	Bovine mRNA fragment for thyroglobulin N-terminal region	3.5
234	D16437	Synechococcus sp. DNA for PacS, complete cds	3.5
235	X97570	Z.mays dek34 gene	3.5
236	S65225	PEP-19=neuron-specific [mice, Genomic, 1358 nt, segment	3.5
		Human immunodeficiency virus type 1 isolate GM4,	
237	U33099	envelope glycoprotein (env) gene, V1-V5 region, partial cds	3.5
238	X71604	H.sapiens son-pseudogene	3.5
239	L47357	Homo sapiens TIMP1 gene, exon 1.	3.5
		Bos taurus immunoglobulin variable region mRNA, partial	
		cds > :: gb AF015492 AF015492 Bos taurus	
240	AF015490	immunoglobulin variable region mRNA, partial cds	3.4
		Arabidopsis thaliana decoy (DECOY) gene, complete cds	
CONTRACTOR OF THE PROPERTY OF	U93308	and valyl tRNA synthetase (valRS) gene, partial cds	3.4
	Y12576	Arabidopsis thaliana mRNA for histone H2B	3.4
243	X07977	Aspergillus amstelodami mtDNA with ARS element	3.4
	X70276	Yeast centromere-containing shuttle vector YCp50	3.4
245	J03268	Yeast (S.cerevisiae) polymerase I gene, complete cds.	3.4
***************************************	AF013168	Homo sapiens hamartin (TSC1) mRNA, complete cds	3.4
*************	M32476	Rat carcinoembryonic antigen-related protein	3.4
248	X83390	Albinaria coerulea complete mithochondria DNA	3.4
249	U63337	Mus musculus cyclin-dependent kinase-2 alpha	3.4
250	D85530	Human CpG island sequence, clone G0310	3.4
		Human FK506-binding protein 12 (FKBP12) mRNA, exons	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
251	M92423	3 and 4, complete FKBP12A mRNA and complete cds.	3.4
252	AB002693	Mouse mRNA for ISBT, complete cds	3.4
253	L01057	Human (clone TRI-6) satellite I repeat region.	3.4
254	U11270	Human antithrombin III gene, exon 1 and partial cds.	3.4
		Rattus norvegicus mRNA for peptide/histidine transporter,	**************************************
255	AB000280	complete cds	3.4
~~~~	J03886	9 No. 1944 A. 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 1865 - 18	3.4
***************************************		Yeast (S.cerevisiae) CLS4 gene encoding a Ca regulatory	
257	M16809	protein, complete cds.	3.4
	X58286	Drosophila genes z600, gdl, Eip28/29 and mex1	3.4
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X07267	Rat gene 33 5'-region	3.4

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALU
*************************	Z68129	H.sapiens IDH gamma gene and TRAP delta gene	3.4
	X87241	H.sapiens mRNA for hFat protein	3.4
262	Z93650	O.bellus 28S rRNA gene, D2 variable region	3.4
263	D87471	Mus musculus mRNA for gsg3, complete cds	3.4
264	L34193	Ipomopsis aggregata chloroplast maturase	3.4
		Sambucus nigra ribosome inactivating protein precursor	
265	AF012899	mRNA, complete cds	3.4
***************************************		Zea mays chloroplast trnC gene, rpoB gene, rpoC1 gene,	<u> </u>
		rpoC2 gene and rps2 gene for transfer RNA-Cys, RNA	
266	X17318	polymerase subunits beta, beta-1, beta-2 and ribosomal	3.4
	L35661	Homo sapiens (subclone H8 4 b9 from P1 35 H5 C8) DNA	3.4
		Human immunodeficiency virus type 1 clone XSH159D1D	
268	U25236	envelope glycoprotein (env) pseudogene, v1-v5 region,	3.4
	L81392	Homo sapiens (subclone 1 c12 from P1 H39) DNA	3.4
	M60172	G.domesticus novel collagen protein mRNA, 3' end.	3.4
۷./۷	171001/2	E.coli araBAD operon encoding L-ribulokinase, L-arabinose	
271	M15263	isomerase, and L-ribulose 5-phosphate 4-epimerase.	3.4
	S53273	OMP=olfactory marker protein {promoter} [mice, Genomic,	and the second s
		antana ana ang managang managa	3.4
213	U22056	Mus musculus fertilin alpha precursor (ADAM 1) mRNA,	3.3
·	******	Antithamnion sp. rhodoplast genes atpI, atpH, atpG, atpF,	
·	X63382	atpD, atpA, orf1, orf2 and orf3	3.3
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U67462	Methanococcus jannaschii section 4 of 150 of the complete	3.3
	M24566	Dictyostelium discoideum tRNA-Glu-GAA gene, clone	3.3
water retorious with our contraction of	L13609	Human catalase (CAT) gene, exon 1, 5' end.	3.3
~~~~~~~	Z11486	Pinus strobus L. mRNA for pine globulin-1	3.3
279	X03366	Bovine spleen trypsin inhibitor II (SI) gene	3.3
		Human DNA for gastric inhibitory polypeptide receptor,	*
280	D49558	exon 5, 6, 7, 8, 9, 10, 11 and 12	3.3
		Sulfolobus solfataricus putative ribokinase and Dbh genes,	***************************************
281	U52110	complete cds, and putative ATPase gene, partial cds	3.3
282	M34663	Human chaperonin (HSP60) non-functional pseudogene 4.	3.3
283	X82303	P.groenlandica mitochondrial cytochrome b gene	3.3
284	D78174	Mouse cerebellum mRNA for Zic4 protein, complete cds	3.3
285	D86966	Human mRNA for KIAA0211 gene, complete cds	3.3
~~~~~~	L13198	Clortridium perfringens type B beta-toxin gene, complete	3.3
***************************************		E.coli leucine-specific transport (LS-BP; LIV-BP) system	····
287	J05516	(livHMGF) genes, complete cds.	3.3
******	M58318	Homo sapiens ala gene.	3.3
	X57297	A. majus TAM1 gene for TNP1 and TNP2	3.3
		Human immunodeficiency virus type 1 isolate GM4,	, _ <del></del>
290	U33099	· · · · · · · · · · · · · · · · · · ·	3.3
270		Coptis japonica mRNA for S-adenosyl-L-	~ • • •
201	D29809		3.3
<b>471</b>	1000	Human T-cell surface antigen T3 delta-chain gene, exons	J.J
202	M12727	2,3,4 and 5, clone pKR-1.	3.3
292	M12727	Human MFD72 dinucleotide repeat DNA > ::	٧.٧
202	V54601	· -	2.2
	X54601	gb I31132 I31132 Sequence 44 from patent US 5582979	3.2
and a comment of the	Y11740	H.sapiens whn gene, exon 1a and 1b	3.2
295	M24566	Dictyostelium discoideum tRNA-Glu-GAA gene, clone	3.2
	0000 50	focal adhesion kinase/pp125FAK/FAK + {alternatively	
296	S83358	spliced} [rats, striatum, mRNA, 4575 nt]	3.2
		Human pulmonary surfactant-associated protein SP-A	
297	M68519	(SFTP1) gene, complete cds.	3.2

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALU
	U22056	Mus musculus fertilin alpha precursor (ADAM 1) mRNA,	3.2
299	AB005803.1	Homo sapiens DNA for histidine-rich glycoprotein,	3.2
300	M24566	Dictyostelium discoideum tRNA-Glu-GAA gene, clone	3.2
301	X66139	M.fascicularis mRNA for epididymal apical protein I	3.2
302	U16955	Plasmodium falciparum ATPase 2 gene, complete cds.	3.2
		Human immunodeficiency virus type 2 (FOPOLC4)	
303	M87108	'polymerase fragment.	3.2
***************************************	U67585	Methanococcus jannaschii section 127 of 150 of the	3.2
*********	4. 16. TH. 1880 A. 1880	Saccharomyces cerevisiae DNA repair/transcription protein	&
305	U70559	Mms19p (MMS19) gene, complete cds	3.2
	D88191	Chlorella Virus vChta-1 gene, complete cds	3.2
	M32352	Mouse renin (Ren-1-d) gene, complete cds.	3.2
	X64406	Marburg Virus RNA for ORF-2 and ORF-3	3.2
200	AU++100	Homo sapiens prospero-related homeobox 1 protein (Prox 1)	
200	NIM 002762 1	mRNA, complete cds	1
and the residence of the second secon	NM_002763.1		3.2
310	Y11681	Homo sapiens mRNA for mitochondrial ribosomal protein	3.2
211	20000	svp[40]=svp-related nuclear receptor/retinoid signaling	
normannen automatika kantan kanta	S80986	modulator [zebrafishes, mRNA, 3876 nt]	3.2
312	Z18630	R.norvegicus mRNA for J1-160/180 neural recognition	3.2
		Sambucus nigra ribosome inactivating protein precursor	
·····	AF012899	mRNA, complete cds	3.2
314	X14336	Filamentous Bacteriophage I2-2 genome	3.1
		Mus musculus SH3-containing protein SH3P7 mRNA,	
315	U58884	complete cds. similar to Human Drebrin	3.1
316	X66139	M.fascicularis mRNA for epididymal apical protein I	3.1
317	D90819	E.coli genomic DNA, Kohara clone #328(39.4-39.8 min.)	3.1
318	U78770	Mus musculus spasmolytic polypeptide (mSP) gene,	3.1
	U06083	Human N-acetylgalactosamine 6-sulphatase	3.1
	THE CONTRACTOR OF THE PROPERTY	Plasmodium falciparum 5.8S ribosomal RNA gene, partial	
		sequence, internal transcribed spacer 2, and large subunit	
320	U48228	ribosomal RNA gene, complete sequence	3
·····	X95188	R.norvegicus mRNA for Pristanoyl-CoA Oxidase	3
		S.scrofa mRNA for protein phosphatase 2A 55 kDa	
322	Z34932	regulatory subunit, alpha isoform (partial)	3
		Ovine adenovirus terminal protein gene, partial cds, 52/55K,	3
		pIIIa, III, pVII, pX, pVI, II, endopeptidase and DNA	
272	U40837		3
omenomorromorromo,comonomorromos		alamanan ara-ara-ara-ara-ara-ara-ara-ara-ara-ara	
***************************************	X77233	Lesculentum (de Ruiter 83G38) Adh2 gene	2.9
~~~~	M81341	Plasmodium falciparum cysteine proteinase gene, complete	2.8
ar cumanom accomenta secue persona de	X99832	H.sapiens CLN3 gene, complete CDS	2.7
321	M64703		2.1
		Homo sapiens DNA for cGMP-binding cGMP-specific	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	AB001635.1	phosphodiesterase (PDE5), exon 21 and complete cds	1.9
minorum manusus menerana nég	X12669	H.sapiens gene for arginase exon 8 and 3'-flanking region	1.8
······································	M19238	Yeast (S.cerevisiae) ribosomal protein L44' gene, complete	1.8
na di Amerika di Karamana mana mana mana mana mana karaman mana karaman mana karaman mana mana mana mana karam	AE001665		1.8
332	X15603	Human elastin gene, exon 1	1.8
333	AE000553.1		1.8
The state of the s	memorani di mandali di dikantan dan kantan di Kantan di Kantan dan dan dan dan dan dan dan dan dan d	obligately oligotrophic bacteria POC-111 DNA for 16S	
334	AB022333	· · · · · · · · · · · · · · · · · · ·	1.8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X51666	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.8
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X16588	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.8
and the second s	U19253	~	1.8

P VALU	DESCRIPTION	ACCESSION	SEQ ID
.7	Haemophilus influenzae Rd section 85 of 163 of the	U32770	***************************************
***************************************	Propithecus verreauxi epsilon globin gene, 5' flanking region		
.7	and exons 1-3, complete cds	U64618	339
.7	Mycoplasma genitalium section 22 of 51 of the complete	U39700	
7		Z82656	
	Homo sapiens mRNA; cDNA DKFZp564P016 (from clone	202030	341
7	<del>-</del>	AT 040227 1	242
. /	;	AL049337.1	342
-	H.sapiens CpG island DNA genomic Mse1 fragment, clone		
.7	36g10, forward read cpg36g10.ft1a	Z60848	and the second s
.7		Z28054	344
	phosphatase inhibitor-2=cytosolic regulatory subunit of type		
.7	1 protein phosphatase [rats, brain, mRNA, 867 nt]	S79213	345
.6	C.anuum mRNA for 1-aminocylopropane-1-carboxylate	X82265	346
	Tetrahymena thermophila polyubiquitin (TTU3) gene,	<u></u>	***************************************
.5	complete cds, and RNA polymerase II subunit 2	U46561	347
.5	Quail fast skeletal muscle troponin I gene, complete cds.	M12132	ANALYSIS CONTRACTOR CONTRACTOR CONTRACTOR
	M.musculus CD22 promoter region	X98097	MATERIAL PROPERTY SECOND SECON
.4	Homo sapiens mRNA for CD151, complete cds	D29963	
	Herpes simplex virus type 2 genomic DNA for 0.74-0.84	D27703	
1.4	region, complete cds	D10471	251
	Micoureus demerarae cytochrome b light strand gene,	D104/1	331
1.3		1124672	252
	mitochondrial gene encoding mitochondrial protein,	U34673	***********************
1.3	Human Pro-tRNA and Leu-tRNA genes.	M15274	
1.3	Trichomonas vaginalis mgl1 gene	AJ000486	354
_	Caenorhabditis elegans cosmid E02H1, complete sequence		
1.3	[Caenorhabditis elegans]	Z47075	355
	Homo sapiens collagen, type XI, alpha 1 type XI collagen		
1.3	(COL11A1) mRNA, complete cds.	NM_001854.1	356
	Clostridium cellulovoran endo-1,4-beta glucanase EngF		porcessors of the water side of the
1.3	(engF) gene, complete cds	U37056	357
1.3	Human cyclin G mRNA, complete cds.	U53328	
	Caenorhabditis elegans cosmid F42G10, complete sequence	······································	
1.3	[Caenorhabditis elegans]	Z48230	359
	Homo sapiens (subclone 2_c7 from P1 H25) DNA sequence.	L42102	
1.3	S. cerevisiae DNA for protein phosphatase T gene	X89417	
	H.sapiens chromosome 22 CpG island DNA genomic Mse1	A09417	301
1.3	, -	770004	262
1.3	fragment, clone 303c5, complete read	Z79884	***
	H.sapiens IDH gamma gene and TRAP delta gene	Z68129	
1.3	Human insulin-like growth factor I (IGF1) gene, intron 4.	U13800	364
	Oryza sativa 3-hydroxy-3-methylglutaryl coenzyme A		
1.3	reductase gene, complete cds.	L28995	
1.3	Saccharomyces cerevisiae Jsn1 gene, complete cds	L43493	366
	Caenorhabditis elegans myo-2 gene for myosin heavy chain		
1.3	2 (MHC-C)	X08066	367
	Clostridium cellulovoran endo-1,4-beta glucanase EngF	***************************************	00x00x00x00x00x00x00x00x00x00x00x0
1.3	(engF) gene, complete cds	U37056	368
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60 E 210 110 110 100 100 10	Clostridium cellulovoran endo-1,4-beta glucanase EngF	U37056 U19905 U74496 U46781 X77300	368 369 370 371

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
6.9 6 (Augusta Arama)		Feline immunodeficiency virus isolate FIV-Pco336-7 pol	***************************************
373	U53745	polyprotein (pol) gene, partial cds	1.3
374	L08266	Mouse Facc mRNA, complete cds.	1.3
375	X52519	Human gene for tyrosine aminotransferase (TAT)	1.3
	***************************************	Aedes aegypti steroid hormone receptor homolog protein	)
376	U87543	gene, complete cds	1.3
		Mycoplasma hyopneumoniae gene for 46 kDa surface	
377	D16682	antigen, complete cds	1.3
378	AF022725	Hordeum vulgare limit dextrinase (HvLD99) gene, complete	1.3
379	L18987	Human alpha-2 type XI collagen mRNA, partial cds.	1.3
380	Y09400	S.scrofa mRNA for apical organic cation transporter protein	1.3
******************************		Human endothelial-monocyte activating polypeptide II	
381	U10117	mRNA, complete cds.	1.3
	U02618	Saccharomyces cerevisiae molasses resistance	1.3
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	
383	Z60848	36g10, forward read cpg36g10.ft1a	1.3
	00.000.0000.000 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400 - 400	H.sapiens gene for pregnancy specific beta-1 glycoprotein >	
***************************************		::: gb S49771 S49771 pregnancy-specific beta 1 glycoprotein	
384	X63203		1.3
	L31854	Mus musculus Ig epsilon-chain C gene, exon 4, M gene,	1.3
	X16588	B. nigra repeat DNA (clone pBN 35)	1.3
····		Xenopus laevis epithelial sodium channel, gamma subunit	
387	U25342	(gammaxENaC) mRNA, complete cds.	1.3
		Neurospora crassa open reading frame gene, complete cds,	
388	L40806	met-10+ gene, complete cds	1.3
·····	S49760	diacylglycerol kinase [rats, brain, mRNA, 3043 nt]	1.3
······································	L15328		1.3
anno anti-	S50809	protein LG=immunoglobulin binding protein Recombinant,	1.3
	ON A STATE OF THE	Baboon herpesvirus HVP2 gB glycoprotein (UL27) gene,	
392	U14662	complete cds.	1.3
		Human intestinal H+/peptide cotransporter (Hpept1) gene,	
393	U13173	complete cds	1.3
	013173	me33e04.r1 Soares mouse embryo NbME13.5 14.5 Mus	1.3
		musculus cDNA clone 389310 5' similar to	
		SW:S105 HUMAN P33763 S100 CALCIUM-BINDING	
394	W71212	PROTEIN A5;. > :: emb X99921 MMS100A13 M.musculus	1 3
J/T	***************************************	Petunia hybrida alcohol dehydrogenase-2 (Adh2) gene,	1.3
305	U25536	promoter, 5'UTR, and partial cds.	1.3
····	X75014		1.3
<del></del>	Z49436	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.3
and the second s	X12780	*************************************	1.3
	X04319	E. coli fhuB gene involved in transport of ferrichrome	1.3
~~~~	U61297	Human progesterone receptor (PGR) gene, far 5' flanking	1.3
400	001297		1.3
401	V00519	Herpesvirus saimiri virion, transformation-associated region, strain C139	1.2
401	X99518	·	1.3
400	N 42 4001	Mink enteritis virus antigenic type 2 capsid protein genes	1.0
402	M24001	·	1.3
400	***	Arabidopsis thaliana geranylgeranyl pyrophosphate synthase	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U44877		1.3
404	AB003431	<i></i>	1.3
		Human steroid 21-hydroxylase (P-450(C21)) B gene,	
-	M12792	complete cds, clone lambda-C21B-1.	1.3

SEQ ID	The state of the s	DESCRIPTION	P VALUE
*****	Y10260	H.sapiens EYA1A gene	1.3
	U41740	Human trans-Golgi p230 mRNA, complete cds	į1.3
409	X02295	Paramecium 5' region of gene for surface protein 51A	1.3
		band 3-related protein=renal anion exchanger AE2 homolog	-
410	S45791	[rabbits, New Zealand White, ileal epithelial cells, mRNA,	1.3
411	AE000625.1	Helicobacter pylori 26695 section 103 of 134 of the	1.3
		R.norvegicus gene encoding alkaline phosphatase, exon 3	
412	X16028	and joined CDS	1.3
413	M73461	Saccharomyces cerevisiae FL100 RNA14 gene, complete	1.3
414	L08845	Drosophila melanogaster disabled mRNA, complete cds	1.3
415	AE000635.1	Helicobacter pylori 26695 section 113 of 134 of the	1.2
416	L39962	Medicago sativa middle repetitive DNA	1.2
THE PROPERTY OF THE PROPERTY O	U55371	Caenorhabditis elegans cosmid T19F4.	1.2
Market Anna Angeles and Anna Anna Anna Anna	X13679	Oryza sativa H3 histone pseudogene H3R-12	1.2
		Homo sapiens epsilon-1 pseudogene (IGHEP1) gene, CH3	- · -
419	J00223	and CH4 regions, exons 3 and 4 and partial sequence	1.2
MINERAL CONTRACTOR CON	AE000649.1	Helicobacter pylori 26695 section 127 of 134 of the	1.2
-	X52256	A.thaliana tufA gene for elongation factor Tu	1.2
		Chilo iridescent virus DNA-directed RNA polymerase and	***
		helicase genes, complete cds's. > :: gb S75674 S75674 DNA	ar weense
		helicase homolog, DNA-depenent RNA polymerase largest	**************************************
422	M81388	· ·	1.2
***************************************	S57565	histamine H2-receptor [rats, Genomic, 1928 nt]	1.2 1.2
*******************************	X84347	H.sapiens mRNA for sperm adhesion molecule hPH-20	1.2
*****	X53579	A.thaliana agamous (AG) gene	1.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Y09539	L.japonicus gene encoding RING finger protein	·
~~~~~	L05500	Human fetal brain adenylyl cyclase mRNA, 3' end.	1.2
~~~~~	X68019	Feline Immunodeficiency Virus GAG gene	1.2
720	AU0017	Human immunodeficiency virus type 2 (HIV-2 ARM)	1.2
		proviral surface glycoprotein (gp125) gene, partial cds. Type	
420	L76739		
***************************************	L38769		1.2
*********************	L39786		1.2
431	L39/80		1.2
423	D10610	Homo sapiens MAT gene for mitochondrial acetoacetyl-	
432	D10510	CoA thiolase, exon 11	1.2
422	1147707	Streptococcus pneumoniae immunoglobulin A1 protease	
433	U47687	(iga) gene, complete cds	1.2
42.4	11/2000	Xenopus laevis beta-transducin repeat containing protein-3	
434	U63922		1.2
43.5	707004	S.tuberosum STACS2 gene for 1-Aminocyclopropane-1-	
***************************************	Z27234	<del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	1.2
436	X65365.1		1.2
	370.400.6	Podospora anserina race A mitochondrial DNA class II	
437	X04336		1.2
	*****	Yeast (S.douglassi) NAM2 gene for mitochondrial leucyl-	
438	X12864	tRNA synthetase (EC 6.1.1.4)	1.2
skiller reprint		Methanosarcina thermophila CO dehydrogenase/acetyl-CoA	
***************************************		synthase alpha subunit (cdhA), epsiolon subunit (cdhB), beta	
concention		subunit (cdhC), and NifH class IV protein homolog genes,	
······································	U66032		1.2
·	L08266	Mouse Facc mRNA, complete cds.	1.2
441	M64085	Mouse spi2 proteinase inhibitor (spi2/eb1) mRNA, 3' end.	1.2
	X12773	karananan kan minaranan mananan nambaran manan man	1.2

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALU
Water and the second se	U13988	Peanut chlorotic streak caulimovirus, complete genome.	1.2
~~~~~	U23180	Caenorhabditis elegans cosmid C28F5	1.2
***************************************	M20537	Mouse thyrotropin beta-subunit gene, exon 5.	1.2
TO STATE OF THE PARTY OF THE PA	U25881	Agrius cingulata NADH dehydrogenase subunit 1 protein,	1.2
***************************************	Y08581	F.rubripes hsp70-4 gene, complete	1.2
	L31848	Homo sapiens serine/threonine kinase receptor 2	1.2
449	M15840	Human interleukin 1-beta (IL1B) gene, complete cds.	1.2
		H. sapiens (D6S443) DNA segment containing (CA) repeat;	
	Z23977	clone AFM277wb5; single read	1.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X14592	P.hybrida chsB gene for chalcone synthase	1.2
452	Z49900	P.sativum mRNA for small GTP-binding protein	1.2
		Haemophilus somnus cryptic prophage genes, capsid	
		scaffolding protein gene, partial cds, major capsid protein	S Williams
		precursor, endonuclease, capsid completion protein, tail	OTTO A SAME AND A SAME
453	U28154	synthesis proteins, holin, and lysozyme genes, complet	1.2
	***************************************	Brassica napus PE3-PEPCase gene for phosphoenolpyruvate	ļ
454	D13987	carboxylase, complete cds	1.2
	D45243	Mouse mRNA for ctk, complete cds	1.2
	***************************************	Caenorhabditis elegans non-alpha nicotinic acetylcholine	} 1 • <del>60</del> {pro-common managements
456	U81144	receptor subunit precursor (unc-29) gene, complete cds	1.2
		Caenorhabditis elegans cosmid H06001, complete sequence	1.2
457	Z92970	[Caenorhabditis elegans]	1.2
management of the second of th	L44118	Homo sapiens proximal CMT1A-REP repeat	&
730	L77110	D.melanogaster achaete gene encoding nerve differentiation,	1.2
450	M17120	complete cds.	1.0
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U55737		1.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X63525	Human ataxia-telangiectasia (ATM) exon 40	1.2
		P.vulgaris loxA gene for lipoxygenase	1.2
and the second s	D16402	Fruitfly Dcdrk gene for Dcdrk kinase, complete cds	1.2
····	M65287	Mouse activin receptor (ActR) mRNA, complete cds.	1.2
404	X13679	Oryza sativa H3 histone pseudogene H3R-12	1.2
4.0	D00404	Aspergillus oryzae pgkA gene for phosphoglycerate kinase,	
465	D28484	complete cds > :: dbj E04898 E04898 gDNA encoding	1.2
Andhuman		Escherichia coli K-12 MG1655 section 173 of 400 of the	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	AE000283	complete genome	1.2
467	X99832		1.2
WANTED TO THE PARTY OF THE PART		Rattus norvegicus follistatin-related protein precursor	
	U06864	mRNA, complete cds.	1.2
469	M87710		1.2
		Human apolipoprotein AI regulatory protein (ARP-1)	
470	M64497	mRNA, complete cds.	1.2
<del></del>	M87710		1.2
····	X60196		1.2
473	X99719	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.2
474	Z95706	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.2
~~~~~~~ <del>~~~~~~~</del>	L76372	·	1.2
		Exogenous mouse mammary tumor virus gene for	
476	D26359	- ,	1.2
		Homo sapiens aldehyde dehydrogenase 7 (ALDH7) mRNA	<b></b>
Women		>:: gb U10868 HSU10868 Human aldehyde dehydrogenase	
477	NM 000694.1		1.2
7//	. 11/1_UUUU74.1		1.2
170	A E015000	Caenorhabditis elegans protein tyrosine phosphatase (ptp-2)	
4/0/	AF015882	mRNA, complete cds	1.2

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID		DESCRIPTION	P VALU
-	U71121	Arabidopsis pyruvate decarboxylase-1 (Pdc1) gene,	1.2
······································	U60804	Danio rerio tumor suppressor p53 (p53) mRNA, complete	1.2
******************	X99254	P.falciparum gene encoding primase, small subunit	1.2
uire adeba, monte con con prop	D10197	Bovine mRNA for histamine H1 receptor, complete cds	1.2
484	Y09764	Homo sapiens GABRE gene, exon 2-8	1.2
		Lycopersicon esculentum class II small heat shock protein	***************************************
tuites in the interest of the contract of the	U72396	Le-HSP17.6 mRNA, complete cds	1.2
Whatevan wassens accesses access	X72950	X.laevis H3l gene for histone H3	1.2
***********	D29956	Human mRNA for KIAA0055 gene, complete cds	1.2
	X56003	E.coli (plasmid pFM205) faeE and faeF genes	1.2
489	M64269	Human mast cell chymase gene, complete cds.	1.2
	4444400000	Human immunodeficiency virus type 1 env gene for	
490	AB002963	envelope glycoprotein, partial cds, clone 205E5B2t	1.2
491	X90846	H.sapiens mRNA for mixed lineage kinase 2	1.2
492	X03715	Spiroplasma melliferum tRNA gene cluster	1.2
		Tropidurus hispidus ATPase subunit 6 (ATPase6) gene,	
493	U83494	mitochondrial gene encoding mitochondrial protein, partial	1.2
494	U60804	Danio rerio tumor suppressor p53 (p53) mRNA, complete	1.2
495	M24081	Tetrahymena pyriformis (clone pTU2) ubiquitin genes, 3'and	1.2
and the state of t		Mus musculus cysteine protease (Lice) gene, exons 3-7, and	
496	U54803	complete cds	1.2
*************	L13748	Human dihydrolipoamide dehydrogenase gene, exon 1.	1.2
*****	U29390	TO THE PROPERTY OF THE PROPERT	1.2
	L81694	on the contract of the contrac	1.2
	X65366.1	R.norvegicus GHF1 gene, intron 2	1.2
***************************************		Human DNA polymerase gamma (polg) gene, promoter	
501	U74651	region and partial cds	1.2
	X65591	S.mansoni mRNA for myosin II heavy chain	1.2
	D17255	Human HepG2 3' region MboI cDNA, clone hmd5c12m3	1.2
*************************	L33792		1.2
		Drosophila melanogaster dead-box protein (dbp45A) gene,	1.2
		complete cds. > :: emb Z23266 DMDEADBXA	
505	L13612	D.melanogaster DEAD-box gene, complete CDS	1.2
***	X81650		1.2
300	A81030	Homo sapiens ankyrin 1, erythrocytic (ANK1) mRNA > ::	1.2
507	NM 000037.1		1.0
descriptions and the second second	X98543	и <del>р Тининания в принентення в постояння принентення постояння в принентення в п</del>	1.2
	D89501		1.2
309	D89301		1.2
510	702174	Human DNA sequence from cosmid B20F6 on chromosome	
CONTRACTOR	Z82174		1.2
511	M36881		1.2
	******	Human clone 121711 defective mariner transposon Hsmar2	
	U92014		1.2
monimum accessors to abuse uses	U09948		1.2
514	M58155		1.2
		Sicilian sandfly fever virus glycoprotein precursor	
~~~~~~~	U30500	\`````````````````````````````````````	1.2
MPADACOCK CONTY OF NEW YORK ON THE	L09190	opo o o o o o o o o o o o o o o o o o o	1.2
***************************************	U67508	· · · · · · · · · · · · · · · · · · ·	1.2
518	M81186	Clostridium botulinum neurotoxin type B (botB) gene,	1.2
		Homo sapiens elastin gene, exons 5-27 and alternatively	
519	U93037		1.2
520	D87454		1.2

Table 2A Nearest Neighbor (BlastN vs. GenBank)

EQ ID	ACCESSION	DESCRIPTION	P VALU
521	D87558	Gallus gallus mRNA for C-Serrate-2, partial cds	1.2
522	Z32847	L.infantum (10541) kinetoplast DNA	1.2
523	U09584	Human PL6 protein (PL6) mRNA, complete cds.	1.2
524	AC001530	Homo sapiens (subclone 2 b8 from P1 H56) DNA sequence	1.2
	X74322	H.sapiens gap-I gene	1.2
~~~~	D29792	Mouse gene for T cell receptor gamma chain	1.2
······································		Mink enteritis virus antigenic type 2 capsid protein genes	ļ
527	M24001	VP1 and VP2, complete cds.	1.2
The same and the same of the s	K02819	Rabbit MHC RLA region class I 19-1 gene, complete cds.	1.2
***************************************	·	Insertion sequence IS1245 (from Mycobacterium avium)	1.2
529	L33879	transposase gene, complete cds.	1.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	AE000607.1	Helicobacter pylori 26695 section 85 of 134 of the complete	
***************************************		me33e04.r1 Soares mouse embryo NbME13.5 14.5 Mus	1.4
		musculus cDNA clone 389310 5' similar to	
	and the same of th	SW:S105 HUMAN P33763 S100 CALCIUM-BINDING	West own
531	W71212	PROTEIN A5;. > :: emb X99921 MMS100A13 M.musculus	1.2
Andrewson of the service of the second	X83078	M.musculus nid gene (exon 4)	1.2
352	1	Rattus norvegicus transcriptional corepressor KAP1/TIF1B	1.2
533	U95041	mRNA, partial cds	1.0
*************************	X58907	H.sapiens CYP21 gene for steroid 21-monooxygenase	1.2
	L11669		1.2
**********	L37053	Human tetracycline transporter-like protein mRNA, Gorilla gorilla (clone Gor-ID) Rhesus-like protein mRNA,	1.2
CONTRACTOR	M33782		1.2
737	17133762	Human TFEB protein mRNA, partial cds.	1.2
520	D70173	Spinacia oleracea mRNA for 26S proteasome beta subunit,	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	D78172		1.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M33782	Human TFEB protein mRNA, partial cds.	1.2
~~~~~	M33782	Human TFEB protein mRNA, partial cds.	1.2
341	Z54312		1.1
5.40	001772	inwardly rectifying K+ channel IRK3(HIT) [hamsters,	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	S81773		1.1
~~~~	Z78910		1.1
544	U08408	Xenopus laevis arginase 3 mRNA, complete cds.	1.1
		Bacillus subtilis 2,3-dihydro-2,3-dihydroxybenzoate	
		dehydrogenase (dhbA), isochorismate synthase (dhbC), 2,3-	
		dihydroxybenzoate-AMP ligase (dhbE), and isochorismatase	
	U26444		1.1
546	X62170		1.1
		Rattus norvegicus proton gated cation channel ASIC1	
547	U94403		1.1
		Homo sapiens DNA for integration site of HBV in a	
	AJ000498	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.1
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X99485		1.1
550	U67520	Methanococcus jannaschii section 62 of 150 of the complete	1.1
annament.		Mycoplasma pulmonis glutamyl tRNA synthetase (gltX)	***************************************
annon		gene, restriction-modification enzyme subunits S1A, R1,	
nonemanna.		M1, S1B (hsds1A, hsdR1, hsdM1, hsds1B) genes, complete	
551	L25415	cds's, DNA polymerase III (polC) gene, 3' region.	1.1
552	X68107		1.1
	X89246	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.1
	X64332	<u> </u>	1.1
		Rattus norvegicus metabotropic glutamate receptor 4b	
555	U47331	·	1.1
·manoromaniana	X16055	ф пошения на поменения на пошения	1.1

SEQ ID	ACCESSION	DESCRIPTION	P VALU
	er general es no, en	Homo sapiens gene for thymidylate synthase, exons 1, 2, 3,	I VALU
557	D00596	4, 5, 6, 7, complete cds	1 1
************	X14639	Tomato ribosomal DNA intergenic spacer	1.1
**********************	U67520	Methanococcus jannaschii section 62 of 150 of the complete	
CONTRACTOR AND	Y11786	R.prowazekii ksgA gene and 2 open reading frames	e Zarone nomen nomen per supremen
	111700	Caenorhabditis elegans cosmid F16C3, complete sequence	1.1
561	Z81065	[Caenorhabditis elegans]	1 1
*********************	X60694	C.perfringens plasmid epsilon-toxin gene	1.1
CONTRACTOR AND	X52648	Schizosaccharomyces pombe p68 gene for p68 protein	1.1
***************************************	X04078	Potato patatin pseudogene (SA10C)	1.1
	204076	Schizosaccharomyces pombe brefeldin A resistance protein	1.1 .
565	U38783		
********************************	U32769	(hba1) and unknown orf genes, complete cds	1.1
management and the second	D89066	Haemophilus influenzae Rd section 84 of 163 of the	1.1
~~~~	U07797	Staphylococcus aureus DNA for DnaA, complete cds	1.1
***************************************	Contract to the contract of th	Rattus norvegicus Sprague-Dawley (T1-alpha) mRNA,	1.1
CONT. POR CONTROL AND ADMINISTRATION AND ADMINISTRA	L14710	C. elegans cosmid K02D10.	1.1
3/0	U42599		1.1
571	1140726	Human epidermal growth factor receptor (EGFR) precursor-	
5/1	U48726	mRNA, exons 8 and 9, partial cds	1.1
570	1120044	Xenopus laevis cyclin-dependent kinase inhibitor p28 gene,	
***************************************	U38844		1.1
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	S82864		1.1
	X65720	M.musculus gene for protein kinase C-gamma (exon1 and	1.1
	D14484	Hepatitis C virus strain J33 genomic RNA, complete genome	1.1
576	L11998	Staphylococcus aureus conjugative transfer gene complex	1.1
		Rice mitochondrion DNA for ATPase subunit 6 and ORFs,	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	D14339	complete cds	1.1
578	D38413		1.1
- Aller Alle		Bacteriophage c-st (from C. botulinum) C1-tox gene for	***************************************
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	D90210		1.1
	X67838	B.napus DNA for myrosinase	1.1
and the second second	X17053	Rat immediate-early serum-responsive JE gene	1.1
582	X12426	Xenopus laevis U1 70K gene exon 4, 5, 6 and 7	1.1
		Bacillus subtilis plasmid pPOD2000 Rep, RapAB, RapA,	
	U55043	ParA, ParB, and ParC genes, complete cds.	1.1
584	M34046	11	1.1
		Clostridium acetobutylicum ATCC 824 F-type ATP synthase	
		subunit a (atpA) gene, F-type ATP synthase subunit c (atpC)	
585	U52367		1.1
586	Z35955	<u></u>	1.1
587	U53179		1.1
		C.herbarum Cla h III mRNA > :: gb I26207 I26207	<u>.</u>
588	X77253		1.1
**********************************	X72713		I.1
MBT-CANCINCALLAND CONCOURS ALARY (THE	U29145		1.1 1.1
		Cubanichthys pengelleyi mitochondrial control region,	1.1
591 T	U06061		1 1
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Market Access Control	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1.1
	**************************************		1.1
and the second s		Mouse oxytocin-neurophysin I gene, complete cds	l.1
J 24 C	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Emericella nidulans septin B (aspB) mRNA, complete cds	1.1
505 N		Human inter-alpha-trypsin inhibitor heavy chain mRNA,	
<b>フラン</b> 1	ATIO123	partial cds, clones lambda-HuHITI-[9,33].	l.1

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
	AND THE PROPERTY OF THE PROPER	immunoglobulin epsilon chain constant region=secreted	ggaggagaagagaagaagaagaagaagaagaagaagaag
597	S53497	form {3' region} [human, B cell myeloma U-266, Genomic,	1.1
		Caenorhabditis elegans calcium channel alpha-1 subunit	
508	U25119	homolog Unc-2 (unc-2) gene, partial cds.	1.1
Market Spring to Commission on the Commission of	U56959	Caenorhabditis elegans cosmid T21F4.	1.1
	X99485	L.luteus mRNA for alpha-subunit of G protein	1.1
	L35848	Homo sapiens IgE receptor beta chain (HTm4) mRNA,	1.1
		Human menin (MEN1) gene, complete cds	1.1
	U93237	Medicago sativa MAP kinase MsERK1 mRNA, complete	1.1
	L07042	N.plumbaginifolia mRNA for catalase (cat3 gene)	1.1
604	Z36977	Rattus norvegicus submaxillary gland alpha-2u globulin	1.1
			1.1
605	J00738	mRNA, complete cds.	I • I
		Xenopus laevis histone gene cluster XlH3-A with genes	
	X03018	H1A, H2B, H3 and H4	1.1
***************************************	X68449	L.esculentum U6 snRNA pseudogene (LeU6.1ps)	1.1
608	U53921	Pneumocystis carinii major surface glycoprotein	1.1
		Human immunodeficiency virus type 2 (FOPOLC2)	300000 V
		polymerase fragment. > :: gb M87107 HIVPOLC3 Human	
609	M87106	immunodeficiency virus type 2 (FOPOLC3) polymerase	1.1
610	U84539	Human dystrobrevin (DTN) gene, exon 11A	1.1
611	J02896	S.purpuratus speract egg protein mRNA, complete cds.	1.1
	J02896	S.purpuratus speract egg protein mRNA, complete cds.	1.1
	AF016253	Klebsiella aerogenes D-amino acid dehydrogenase	1.1
		Saccharomyces cerevisiae aminonitrophenyl propanediol	<u> </u>
		(ANP1), UV excision repair protein (RAD23), cytochrome c	
614	L22173	isozyme (CYC7) genes, complete cds.	1.1
	U41357	Tetrahymena thermophila P-type ATPase (TPA11) gene,	1.1
	X14383	Bunyamwera virus L protein RNA, complete cds	1.1
	71700	Mus musculus DNA repair enzyme (Ku 70) gene, exon 13	
617	U50378	and complete cds	1.1
U17	030376	Mus musculus of 5HT5 receptor cDNA gene encoding	
610	Z18278	5HT5 serotonin receptor	1.1
	U14172	Mus musculus p162 protein mRNA, complete cds.	1.1
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		1.1
620	X76762	Bacillus subtilis CzcD (czcD) gene, partial cds, TrkA (trkA)	1.1
<b>601</b>	1100055		1.1
which had been the second entering an expension as per	U62055	Igene, complete cds	1.1
	X81847	E. caroto (ora pora, pora man-pora garante	
	M25477	Caenorhabditis elegans collagen (col6) gene, complete cds.	1.1
	X97196	D.melanogaster X gene	1.1
	L39059	Homo sapiens transcription factor SL1 mRNA, complete	1.1
626	M21790	X.laevis complement component 3 (C3) mRNA, 3' end.	1.1
	·	H.sapiens brca2 gene exon 11 > :: emb A62786.1 A62786	NEW TOWN
	X95161	Sequence 27 from Patent WO9719110	1.1
628	K02446	Chicken smooth-muscle alpha-tropomyosin gene, complete	1.1
629	U24171		1.1
630	Z26306	H.sapiens isoform 1 gene for L-type calcium channel, exons	1.1
631	U23070	Human putative transmembrane protein (nma) mRNA,	1.1
	AE000046	Mycoplasma pneumoniae section 46 of 63 of the complete	1.1
	X17548	D. melanogaster singed gene, exon 1	1.1
	U54497	Xenopus laevis integrin alpha 4 mRNA, complete cds.	1.1
	J02793	Mouse R and L1 (3' end) repetitive elements.	1.1
*********	D10832	Equine herpesvirus genomic DNA for 'TREC' OCT-	1.1
	X51766	L.polyphyllus mRNA for ribosomal protein S16	1.1

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
638	U60804	Danio rerio tumor suppressor p53 (p53) mRNA, complete	1.1
639	Z79068	H.sapiens flow-sorted chromosome 6 HindIII fragment,	1.1
640	Z73585	S.cerevisiae chromosome XVI reading frame ORF	1.1
***************************************		Cricetulus griseus SRD-2 mutant sterol regulatory element	
641	U22818	binding protein-2 (SREBP-2) mRNA, complete cds.	1.1
642	M25710	Human thyroid peroxidase (TPO) gene, exon 11.	1.1
643	L12591	Human alkaline phosphatase gene, 5' flanking region.	1.1
644	L25616	Homo sapiens kinectin mRNA, complete cds	1.1
ALLONDONOS DA POR PORTA	and the second s	Bos taurus clone 9 immunoglobulin lambda light chain	
645	U32251	variable region (Vlambda1b) mRNA, partial cds	1.1
	M97516	Mouse alpha-2 adrenergic receptor gene, complete cds.	1.1
	AJ000060	Mus musculus gene encoding lysosomal hayluronidase,	1.1
	113000000000000000000000000000000000000	Simian varicella virus tegument protein gene, complete cds;	
		serine/threonine protein kinase genes, complete cds;	
618	L07067	membrane glycoprotein genes, complete cds's.	1.1
and the second s	L00619	Mouse T-cell-specific tyrosine kinase (Itk) mRNA, complete	5 (~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
AND THE PROPERTY OF THE PARTY O	M76981	Glycine max vspA gene, complete cds.	1.1
	\$	Human mRNA for KIAA0270 gene, partial cds	1.1
	D87460		Ç
	D17466	Streptomyces setonii crtS gene for sigma factor, complete	1.1
	D89066	Staphylococcus aureus DNA for DnaA, complete cds	1.1
654	X96853	; - <b>P</b>	1.1
		Homo sapiens potassium intermediate/small conductance	on digital control of the control of
		calcium-activated channel, subfamily N, member 1	
	00000000	(KCNN1) mRNA > :: gb U69883 HSU69883 Human	
655	NM_002248.1	calcium-activated potassium channel hSK1 (SK) mRNA,	1.1
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	AND ADDRESS AND AD
656	Z63493	85c1, reverse read cpg85c1.rt1a	1.1
657	M21995	Human coagulation factor XIII a subunit gene, exon 9.	1.1
658	X60367	Mouse CRBPI mRNA for cellular retinol binding protein I	1.1
659	M63224	Wheat germin 9f-3.8 gene, complete cds.	1.1
	<u></u>	Mouse GA binding protein (GABP-alpha subunit) mRNA,	
660	M74515	complete cds.	1.1
	X52949.1	G.intestinalis DNA for rRNA tandem repeat unit	1.1
	·	Rattus norvegicus gene for ribosomal protein S15, exon 1, 2,	
662	D11388	3, 4, complete cds	1.1
	U22302	Human histo blood group ABO glycosyltransferase	1.1
	M38132	S.pombe rad1 gene, complete cds.	1
007	14130132	Perissodus microlepis T51a mitochondrion NADH	
665	1107261	dehydrogenase subunit 2 gene, complete cds	1
	U07261	S.cerevisiae chromosome XV reading frame ORF YOR336w	\$
A-40-00-404-2007-00-00-00-00-00-00-00-00-00-00-00-00-	Z75244		1
00/	D63816	Mouse DNA for glutamate transporter Slc1a3, exon 1	1
	T-10.4.600	Euproctus platycephalus mitochondrion 16S rRNA gene,	
	U04699	partial sequence.	1
CONTRACTOR	M19881	P.falciparum knop protein (KP) gene, complete cds.	1
	X64310	H.sapiens DNA for pu-py sequence on 11p13	1
671	X56469	Mouse HSA-A gene coding for heat stable antigen	1
	***************************************	Meloidogyne hapla mitochondrial COII gene, 3' end of cds;	
	nonimonoral	transfer RNA-His gene; 16S ribosomal RNA gene; ND3	
672	L76262	gene, complete cds; cytochrome b (cytb) gene, 5' end of cds.	1
	X52574	Mouse mRNA from Mov10 locus	1
***************************************	Z11711	H.sapiens gene for alpha-2 macroglobulin, exon 1	1
	M37240	P.juncea N8 family repetitive sequence DNA.	1
****	Z72947	S.cerevisiae chromosome VII reading frame ORF YGR162w	` 1

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
677	X58713	N.crassa phr gene for photolyase	1
		Human immunodeficiency virus type 2 (HIV-2), complete	
678	M30502	proviral genome.	1
679	M23082	Chicken embryo fibroblast tropomyosin mRNA, complete	1
680	K03203	Human PRH1 locus salivary proline-rich protein mRNA	1
681	M86844	Microtus chrotorrhinus tandem satellite array DNA sequence	1
682	U55381	Leishmania tropica P-glycoprotein E gene, complete cds	1
		Rattus norvegicus gene for ribosomal protein S15, exon 1, 2,	
683	D11388	3, 4, complete cds	1
684	L43496	Xenopus laevis ligase I (ligI) mRNA, complete cds	1
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X69103	C.glutamicum csp2 gene	0.99
~~~~~	M32883	Alfalfa leghemoglobin gene, complete cds.	0.98
		Human immunodeficiency virus type 2 (HIV-2), complete	
687	M30502	proviral genome.	0.98
	K02212	Human alpha-1-antitrypsin gene (S variant), complete cds.	0.96
AND	Y09746	H.oligactis mRNA for heat shock protein 70	0.96
SANGER OF THE OWNER	D12580	Group II phytoplasma gene for 16S ribosomal RNA	0.95
	***************************************	Haematobia irritans (clone Horn.fly.3.7) mariner transposase	<del></del>
691	L10465	pseudogene, partial cds.	0.95
	U22541	Enterococcus hirae insertion sequence.	0.95
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Caenorhabditis elegans multidrug resistance related protein 2	&,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
693	U66261	(mrp-2) mRNA, complete cds	0.93
***********	L05517	Plasmodium chabaudi DNA sequence, exon 1.	0.91
021		NSCL-2=basic domain helix-loop-helix gene [mice, embryo,	V.7.1
695	S40532	mRNA, 2230 nt]	0.91
		Pseudomonas aureofaciens phzFABCD genes, complete	
696	L48339	cds's. > :: gb AR008980 AR008980 Sequence 11 from patent	0.65
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X77515	R.rubrum nifJ gene	0.65
		Xenopus mRNA for APEG protein, containing a highly	
698	X51394	repetitive amino acid sequence	0.65
~~~~~~	AF097906	Rana catesbeiana myosin heavy chain (MHC-3) mRNA,	0.64
arren manerana arren	X64724	M.musculus NKR-P1 2 gene for natural killer cell receptor,	0.62
		Homo sapiens formyl peptide receptor-like 1 (FPRL1)	
		mRNA > :: gb M84562 HUMFPRL1A Human formyl	
701	NM 001462.1	peptide receptor-like receptor (FPRL1) mRNA, complete	0.61
	AF110966.1	HIV-1 isolate C-96BW04.10 country Botswana, complete	0.6
	Y10743	active mark commonwear and accommon acc	0.6
		Xenopus laevis pyruvate dehydrogenase E1-beta subunit	<u> </u>
704	U39669	(PdhE1beta-2) mRNA, partial cds	0.59
and the second s	AJ004952		0.59
	AF125454		0.59
***************************************	X59002	Leukemogenic retrovirus T1223/B env gene	0.59
	Z73039	S.cerevisiae chromosome VII reading frame ORF YGR254w	
	Z28236		0.58
		Xenopus laevis pyruvate dehydrogenase E1-beta subunit	
710	U39669	(PdhE1beta-2) mRNA, partial cds	0.58
	Z16651	H. sapiens (D10S199) DNA segment containing	0.58
Contract of the Contract of th	X60163	enganananananananananananananananananana	0.58
114	/xUV1UJ	Zymomonas mobilis invA gene for intracellular invertase	v.20
712	D10465	E1, complete cds	0.58
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z33280	M.capricolum DNA for CONTIG MC376	0.58
	LJJ40U	1vi.capitotuiti DNA 10t CONTTO IviC3/0	V.J/
,17		Sambucus nigra ribosome inactivating protein precursor	

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
716	AB014572	Homo sapiens mRNA for KIAA0672 protein, complete cds	0.57
		Homo sapiens protein phosphatase 1, regulatory subunit 10	
717	NM_002714.1	(PPP1R10) mRNA > :: emb Y13247 HSFB19 Homo sapiens	0.57
718	M10630	Bacteriophage U3 cleavage site for phage phi-X174 gene A	0.56
umanuman <del>an anta diferen</del> t	THE RESIDENCE OF THE PROPERTY	H.sapiens CpG island DNA genomic Mse1 fragment, clone	
719	Z55474	42d3, reverse read cpg42d3.rt1a	0.56
720	U96697	Drosophila melanogaster DPP2C1 (dpp2c1) mRNA,	0.56
721	AC001048	Homo sapiens (subclone 1 f12 from P1 H55) DNA	0.56
722	M32245	Human aromatase gene, 5' flank.	0.55
723	AF051944	Gallus gallus Xin mRNA, complete cds	0.55
anno anti-anti-anti-anti-anti-anti-anti-anti-	AF077539	Caenorhabditis elegans cosmid T25D3	0.54
		Entamoeba histolytica U6 small nuclear RNA gene,	<u> </u>
725	U43841	complete sequence	0.54
	U-130-11	Homo sapiens von Hippel-Lindau syndrome (VHL) mRNA,	
726	NM 000551.1	and translated products	0.54
120	14141_000551.1	Cavia porcellus interleukin-5 receptor alpha precursor (gpIL-	
727	U55215	Sra) mRNA, complete cds	0.53
	·	Human mRNA, Xq terminal portion	0.53
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	D16471	S.cerevisiae NOP77 gene for essential nucleolar protein	0.52
arana arang ar	X76245		0.52
<del>varoninamente de la constanta /del>	Z11993	V.proteolyticus aminopeptidase	0.49
	L34620	Eubacterium fosser 16S ribosomal RNA.	0.49
	U28757	Sus scrofa lysozyme gene, complete cds	
733	U75187	Arabidopsis thaliana germin-like protein (GLP1) mRNA,	0.49
		H.sapiens telomeric DNA sequence, clone 2PTEL005, read	0.40
	Z96514	2PTELOO005.seq	0.48
	AE000579.1	Helicobacter pylori 26695 section 57 of 134 of the complete	
	Y13852		0.47
CONTRACTOR	X14036	Tomato cab-7 gene for type II chlorophyll a/b-binding	0.47
	X84308		0.47
739	S78378	Oct-4 (t12 haplotype) [mice, Genomic, 1191 nt, segment 3	0.46
		Homo sapiens 5-methyltetrahydrofolate-homocysteine	
740	NM_000254.1	methyltransferase synthase mRNA, complete cds	0.46
		Plasmodium falciparum (clone Pfg377 [PfsXLX]) DNA	
741	L04161	sequence, complete cds	0.46
742	X95887	H.sapiens PLP gene, intron 1	0.45
***************************************		Rattus norvegicus cyclic nucleotide phosphodiesterase	
743	U38179	(RNPDE3A) mRNA, complete cds.	0.45
***************************************		Macaca mulatta MHC class I antigen Mamu B*02 mRNA,	
744	U41833	complete cds	0.45
~~~~~~~~~	90.000.000.000.000.000.000.000.000.000.	Plasmid pAD1 (from Enterococcus faecalis) replication-	
745	L01794	associate protein (repA, repB, and repC) genes, complete	0.45
	······································	Human connexin43 gap junction protein (connexin43) gene,	
746	U64573	exon 1 and promoter region	0.44
************************	X60832	P.sativum gene (GS 3A) for glutamine synthase	0.44
anno altra compressione di compressioni di construiri di	Z93997	Unidentified bacterium DNA for 16S ribosomal RNA	0.44
CONTRACTOR OF THE PROPERTY OF	U32818	Haemophilus influenzae Rd section 133 of 163 of the	0.44
		Sphaerozoum punctatum 16S-like ribosomal RNA gene,	ļ
750	AF018161	complete sequence	0.44
	D78156	Human mRNA for rasGTPase activating protein, partial cds	0.44
~~~~~	AB000173	Porcine mRNA for endopeptidase 24.16, complete cds	0.44
	M36626	Rat simple sequence DNA, clone 5.	0.44
133	1/120070	M.musculus flanking region of exon 1 of SEZ-6 gene	V.TT
		IVI.HUSCUIUS HAIRKIIK IEKIOH OI CAUH I OI SEZ-O KUIC	1

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
		Caldocellum saccharolyticum celB gene for	***************************************
	X13602	cellobiohydrolase/endocellulase	0.44
756	AF005664	Homo sapiens properdin (PFC) gene, complete cds	0.44
		Chinese hamster cAMP-dependent protein kinase, catalytic	
757	M63312	subunit-beta mRNA, complete cds.	0.44
758	U43382	Human Down Syndrome region of chromosome 21 DNA.	0.44
759	U75187	Arabidopsis thaliana germin-like protein (GLP1) mRNA,	0.44
Marital Add Add Add Ada San San San San San San San San San Sa		Human duplicate spinal muscular atrophy mRNA, clone	<u> </u>
760	U21914	5G7, partial cds.	0.43
**********	AF010537	Plasmodium falciparum microsatellite TA3 sequence	0.43
762	X05034	Rat C2A gene for prostatic binding protein (PBP)	0.43
metacrarecoccaciones accessos acces	·	Dictyostelium discoideum mitochondrial DNA for NADH	
763	D16579	dehydrogenase subunits and cytochrome oxidase subunit	0.43
	M58155	African swine fever virus multigene families 360 and 110.	0.43
	Z82587	R.prowazekii genomic DNA fragment (clone A315R)	0.43
	202501	Bacillus subtilis DNA for glyceraldehyde-3-phosphate	0.43
766	X13011	dehydrogenase (EC 1.2.1.12)	0.42
<del>~~~~</del>	X59952	T.thermophila SB2040 micronuclear limited DNA element	0.43
	Z70730		0.43
	X94445	L.lactis gene for beta-phosphoglucomutase	0.43
and the second s	WATER MANAGEMENT WOODS TO THE TOTAL PROPERTY OF THE PROPERTY O	S.pombe cwl1 gene	0.43
170	X63628	S.pombe MFm2 gene	0.43
	******	O.berteriana mitochondrial nad5 gene for NADH	volue. August 1
	X60049	dehydrogenase subunit 5, exons d and e	0.43
	D45241	Rat MHC class II gene (RT1.DOa), exon 2, 3, 4 and 5	0.43
····	D83472.1	Bos taurus gene for adrenodoxin reductase, exon 1, 2	0.43
774	M34044	Pig Na+/glucose cotransporter protein (SGLT1) mRNA, 3'	0.43
		Human putative G protein-coupled receptor (AZ3B) mRNA,	
more commence and a second	U28488	complete cds	0.43
	X89398	H.sapiens ung gene for uracil DNA-glycosylase	0.43
***************	Z36879		0.43
····	M73760	Mouse serine protienase, complete cds.	0.43
untermountaines en marie en participar que pièc	Z36803	H.sapiens (xs151) mRNA, 355bp	0.43
~~~~~~~~~~~	M33940	Human Val-tRNA and Lys-tRNA genes.	0.43
781	M96159	Rattus norvegicus adenylyl cyclase type V mRNA, complete	0.43
782	X54134	Human HPTP epsilon mRNA for protein tyrosine	0.43
783	Z73039	S.cerevisiae chromosome VII reading frame ORF YGR254w	0.43
784	AB002312	Human mRNA for KIAA0314 gene, partial cds	0.43
785	AB002312	Human mRNA for KIAA0314 gene, partial cds	0.43
		Human ATP binding cassette transporter (ABCR) mRNA,	
786	U88667	complete cds	0.43
***************************************	kin kinin menenen menen menen menen kenen kan menen kan menen pen berangan pen berangan pendangan dan sebagai	Human ATP binding cassette transporter (ABCR) mRNA,	
787	U88667	complete cds	0.43
***************************************		H.sapiens CpG island DNA genomic Mse1 fragment, clone	J.13
788	Z65552	46b12, reverse read cpg46b12.rt1a	0.43
	X15599	Phanerochaete chrysosporium LIP2 gene for lignin	0.43
		Drosophila melanogaster dopamine D1-like receptor mRNA,	V.TJ
	U22106	partial cds.	0.42
790	U	Mouse transforming growth factor beta mRNA	0.43 0.43
occourant announcement for a	M13177		1144
occourant announcement for a	M13177		······
occourant announcement for a	M13177	Homo sapiens hormone receptor (growth factor-inducible	······································
791	<b>NOTHER MORNING</b> (CO. M. CO. M	Homo sapiens hormone receptor (growth factor-inducible nuclear protein N10) (HMR) mRNA > ::	protection of the letter of headenstands concerns
791	NM_002135.1	Homo sapiens hormone receptor (growth factor-inducible	protection of the letter of headenstands concerns

SEQ ID	ACCESSION	DESCRIPTION	P VALU
794	M60858	Human nucleolin gene, complete cds.	0.42
		Caenorhabditis elegans cosmid F07D3, complete sequence	<u> </u>
795	Z71179	[Caenorhabditis elegans]	0.42
796	Z11804	D.discoideum ras gene	0.42
	**************************************	Equus caballus (clone GEN2-9) germline Ig lambda chain	***************************************
		gene, J4-C4 region, last 2 exons. > :: gb L07573 HRSIGLL	8000 MAR
797	L07572	Equus caballus (clone HVL1) germline Ig lambda chain	0.42
**************************************	L25637	Xenopus laevis HNF-3beta gene, complete cds.	0.42
********************************	000 Mar-186 Sum Antonio Maria III (1974) (1974) (1974) (1974) (1974)	Escherichia coli K-12 MG1655 section 183 of 400 of the	
799	AE000293	complete genome	0.42
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	
800	Z55829	68a3, forward read cpg68a3.ft1a	0.42
WENTER PROPERTY OF THE PROPERT	Y11280	A.vinelandii yoh-1, ibpB, cynR, leuC, leuD, leuB, asd, usg-1	
*****	Y08925		·÷····
	X89961	P.falciparum aarp3 gene, exon	0.42
***************************************	X82330	H.sapiens DNA for MCS gene	0.42
<b>8</b> 04	A8233U	A.hypogaea chi2.2 gene for chitinase (class II)	0.42
905	D00655	Rattus norvegicus mRNA for scavenger receptor class B,	
and the second s	D89655	complete cds	0.42
806	U70730	Human SnoN2 mRNA, complete cds	0.42
		M.musculus flanking region of exon 1 of SEZ-6 gene	-
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Y09922	including promoter sequence	0.42
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	X57638	Mouse mRNA for peroxisome proliferator activated receptor	ć
809	M33196	Human Fc-epsilon-receptor gamma-chain gene, complete	0.42
		Caldocellum saccharolyticum celB gene for	
810	X13602	cellobiohydrolase/endocellulase	0.42
		Caldocellum saccharolyticum celB gene for	
811	X13602	cellobiohydrolase/endocellulase	0.42
and the second		Mycoplasma pulmonis putative lipoprotein (lipA), VsaB	1
, company		lipoprotein (vsaB), VsaC2 lipoprotein (vsaC2), VsaE2	
Proportion Annabe		lipoprotein (vsaE2), VsaD lipoprotein (vsaD) genes, partial	Tona Carlo
812	U23947	cds, VsaA lipoprotein (vsaA) gene, complete cds, VsaC1 l	0.42
www.www		Sus scrofa clone ARO34B cytochrome P450 aromatase	
813	U37312	mRNA, complete cds	0.42
		Phanerochaete chrysosporium manganese peroxidase	***************************************
814	U70998	isozyme 3 (mnp3) gene, complete cds	0.42
	**************************************	Phanerochaete chrysosporium manganese peroxidase	·····
815	U70998	isozyme 3 (mnp3) gene, complete cds	0.42
	X59379	Mouse mRNA for amyloid beta precursor (protease nexin II)	
varance construction and a state of the	X54134	Human HPTP epsilon mRNA for protein tyrosine	0.42
verrenen er	Z32676	H.sapiens x11 gene, promoter region	0.42
		Rattus norvegicus protein tyrosine phosphatase D30 mRNA,	···-
819	U28938	i r	0.42
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	L41867		0.42
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	AF004659		0.42
ranconario en encomo conocidado	U75187	<del>gan talan and an </del>	77.10907874C.750.45040.46040.46040404040404040404
522	~ , J	Chlamydomonas reinhardtii ADP-ribosylation factor (ARF)	0.42
872	U27120		0.42
043	UZ/1ZU		0.42
sonna wood		Chlamydia psittaci RecJ recombination protein gene, partial	
WWW PETERON		cds and ORF2, ORF4, glutamyl-tRNA synthetase, outer	
004		membrane protein 3 (omp3), outer membrane protein 2	
824	U <b>41</b> 759		0.41
	0.1051.6	Candida maltosa ALK4 (CYP52D1) gene for n-alkane	
825]	D12716	inducible cytochrome P-450, complete cds	0.41

EQ II	tioned to the control of the control	DESCRIPTION	P VALI
	6 J01261	P.vulgaris lectin gene, complete cds.	0.41
***	7 K00131	mouse b2 repeat sequence from clone mm61.	0.41
	8 AF000949	Canis familiaris keratin (KRT9) gene, complete cds	0.41
decreases or the property of	9 U67580	Methanococcus jannaschii section 122 of 150 of the	0.41
William Arrage arrays	0 X98568	H.sapiens type X collagen gene	0.41
······································	1 X13595	Bean DNA for glycine-rich cell wall protein GRP 1.0	0.41
	2 Y13852	Drosophila diplacantha clone D3 inactive Bari-1 family	0.41
83	3 X53815	Human G6PD gene for glucose-6-phosphate dehydrogenase. Bovine (clone p59K2) 3',5'-cyclic nucleotide	0.41
	Redictables of a regulation	phosphodiesterase (BTPDE1A1) mRNA, complete cds. > :: gb I30433 I30433 Sequence 16 from patent US 5580771 > ::	SOMEONIA (SOC
83	4 L34069	gb I35677 I35677 Sequence 16 from patent US 5602019 > ::	0.41
83.	5 U26425	Human phospholipase C-beta-3 (PLCB3) gene, complete	0.41
		Homo sapiens transforming growth factor-beta 1	
83	6 J04431	gb I14074 I14074 Sequence 1 from patent US 5445941	0.41
83′	7 U32802	Haemophilus influenzae Rd section 117 of 163 of the	0.41
838	8 U24676	Drosophila melanogaster twinstar (tsr) gene, complete cds	0.41
***********************		Bacillus thuringiensis kurstaki insecticidal delta-endotoxin	0.41
839	U87793	CryIA(c) (cryIA(c)) gene, complete cds	0.41
		Mus musculus domesticus insulin-like growth factor 2	0.41
840	L06446	receptor (Igf2r) gene, exon 2 and partial cds	0.41
	7,200-7-0	Candida transcalia POVO cono formanio cos	0.41
841	X06660	Candida tropicalis POX9 gene for peroxisomal catalase	
*****************	2 U05349	(PXP-9) > :: dbj E01922 E01922 DNA encoding catalase of	0.41
042	1003349	Equine rotavirus glycoprotein VP7 mRNA, complete cds.	0.41
0.40	L12058	Clostridium sordellii 7-alpha-hydroxysteroid dehydrogenase	
****	X71327	gene, complete cds.	0.41
	AE000534.1	M. musculus mRNA for MRE-binding transcription factor	0.41
04.	AEUUU334.I	Helicobacter pylori 26695 section 12 of 134 of the complete	0.41
916	L33727	Mouse 43kDa acetylcholine receptor-associated protein	The second secon
**********	Agreement and the contract of	(Rapsn) gene, exons 3-8	0.41
04/	Y11270	E.histolytica INO1 gene	0.41
0.40	T 10200	Camptotheca acuminata 3-hydroxy-3-methylglutaryl	and the same of th
	L10390	coenzyme A reductase gene, complete cds.	0.41
******************	M34044	Pig Na+/glucose cotransporter protein (SGLT1) mRNA, 3'	0.41
·····	M20543	Human skeletal alpha-actin gene, complete cds.	0.41
851	U97192	Caenorhabditis elegans cosmid C01F4	0.41
o = -	****	Mus musculus strain BALB/c IgH C alpha gene, 3'	
************	U65625	regulatory region enhancer E	0.41
***************************************	X78344	S.cerevisiae CAT8 gene	0.41
854	Z26280	H.sapiens isoform 1 gene for L-type calcium channel, exon	0.41
855	Z23995	H. sapiens (D1S469) DNA segment containing (CA) repeat; clone AFM280za5; single read	0.41
~ -		Xenopus mRNA for APEG protein, containing a highly	
Pina footpateron voge on structure year			0.41
***************************************	U72719		0.41
858	M95673	C 1	0.41
		Rattus norvegicus protein tyrosine phosphatase D30 mRNA,	
859	U28938		0.41
		Homo sapiens UDP glycosyltransferase 1 phenol UDP-	·····
860	NM 001072.1	1 1 2	0.41
Marine Ma	~~~~~~ <del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	S.cerevisiae chromosome VII reading frame ORF YGR099w	0.71 0.41
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	77	************************************
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	***************************************	Human myoglobin gene, exon 2.	0.4

EQ ID	ACCESSION	DESCRIPTION	P VALUI
864	D00844	Gallus gallus mRNA for virus activating protease, complete	0.4
		Homo sapiens mRNA for 6-phosphofructo-2-kinase/fructose	J
865	D49817	2, 6-bisphosphatase, complete cds	0.4
866	L78742	Homo sapiens (subclone 10 b11 from P1 H16) DNA	0.4
867	X51772	Rat DNA for calpain II large subunit (EC 3.4.22.17) (exon	0.4
***************************************		Luxilus chrysocephalus chrysocephalus Meramec R.,	
		Jefferson Co., MO, USA, mitochondrion 12S rRNA and 16S	Season of
868	U09473	rRNA genes, partial sequence, and tRNA-Val gene,	0.4
869	X52614	Maize chloroplast rps15 gene for ribosomal protein S15	0.4
***************************************		Sambucus nigra ribosome inactivating protein precursor	U. 7
870	U76524	mRNA, complete cds	0.4
*****************	<del></del>	D.discoideum uridine diphosphoglucose pyrophosphorylase	10.4
871	M30467	(UDPGP1) gene, 5' end.	0.4
		Homo sapiens potassium intermediate/small conductance	0.4
	90000000	calcium-activated channel, subfamily N, member 1	
	***************************************	(KCNN1) mRNA > :: gb U69883 HSU69883 Human	
872	NM 002248.1	calcium-activated potassium channel hSK1 (SK) mRNA,	0.4
WWW.	M81686	Petunia hybrida Sx-protein	0.4
******	S39508	myelin P2 [mice, Genomic, 3694 nt]	0.4
*****	U82966	Oryza sativa Ca2+-ATPase gene, complete cds	0.4
*******	X16323	Human mDNA for hands and the first (HOE)	0.4
070	A10323	Human mRNA for hepatocyte growth factor (HGF)	0.4
877	U31631	Dictyostelium discoideum class II apurinic/apyrimidinic(AP)	1
~~~~~	X60753	endonuclease (DdapeA) gene, complete cds	0.4
0/0	AUU/33	P.sylvestris BBS gene for bibenzyl synthase	0.4
870	V00711	Mus musculus mitochondrial genome > ::	•
017	V 00/11	gb J01420 MUSMTCG Mouse mitochondrion, complete	0.4
ያያለ	U66274	Rattus norvegicus neuropeptide Y5 receptor (NPYR5)	
or the second contract of the second	X83673	mRNA, complete cds	0.4
~~~~	X14036	X.laevis SSB1 gene	0.4
002	A14030	Tomato cab-7 gene for type II chlorophyll a/b-binding	0.4
002	1176524	Sambucus nigra ribosome inactivating protein precursor	
	U76524 X64467	mRNA, complete cds	0.4
····	······································	H.sapiens ALAD gene for porphobilinogen synthase	0.4
000	L81673	Homo sapiens (subclone 1_f8 from P1 H49) DNA sequence	0.4
996	1 2/007	Saccharomyces cerevisiae mitochondrion transfer RNA-	
	L36887	Leu, Gln, Lys, Arg, Gly, Asp, Ser2, Arg2, Ala, Ile, Tyr, Asn	0.4
<del>~~~~</del>	L24905	Bombyx mori nuclear polyhedrosis virus hr5 gene.	0.4
······································	K00841		0.4
	K02819		0.4
******	L24905		0.4
	X69491		0.4
892	X95887		0.4
		H.sapiens telomeric DNA sequence, clone 3PTEL001, read	
893	Z96576		0.4
		H. sapiens (D1S196) DNA segment containing (CA) repeat;	ALINE AND THE TOTAL AND
	Z16503		0.4
895]	M20572	Mouse interleukin 6 (IL-6) gene, complete cds.	0.4
ana mana ang ang agai	AF022372		0.4
897	Z77964	The state of the s	0.4
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	***************************************
898	Z58874		0.4
	X53422	D. grimshawi s18, s15, s19 and s16 chorion protein genes	

Table 2A Nearest Neighbor (BlastN vs. GenBank)

EQ ID	ACCESSION	DESCRIPTION	P VALU	
	**************************************	Homo sapiens mannose phosphate isomerase (MPI) mRNA		
		> :: emb X76057 HSRPMI H.sapiens PMI1 mRNA for	PROFESSION AND ADDRESS OF THE PROFES	
900	NM_002435.1	phosphomannose isomerase	0.4	
		D. virilis mitochondrial DNA for origin of replication, small	economy o	
		mitochondrial ribosomal RNA, transfer tRNAs tRNA-fMet,		
************	X05914	tRNA-Gln and tRNA-Ile	0.4	
******************	Z95179	G.gallus microsatellite DNA (LEI0290 (= EC11EO7))	0.4	
903	X65627	M.musculus mRNA TNZ2 for p68 RNA helicase	0.4	
		Salmonella typhimurium hydroxyethyl thiazole kinase (thiM)	)	
904	U87940	and HMP-P kinase (thiD) genes, complete cds	0.4	
	0000	Sus scrofa clone pvg13 Ig heavy chain variable VDJ region		
905	U15450	mRNA, partial cds.	0.4	
		Rat cardiac specific sodium channel alpha-subunit mRNA,		
************************	M27902	complete cds.	0.4	
907	U22893	Rattus norvegicus muscle Y-box protein YB2 mRNA,	0.4	
	50-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-	Homo sapiens neurexin 4 (contactin associated protein)		
		(NRXN4) mRNA > :: gb U87223 HSU87223 Homo sapiens		
908	NM_003632.1	contactin associated protein (Caspr) mRNA, complete cds	0.4	
		Homo sapiens myasthenia gravis autoantigen gravin mRNA,		
Description of the second	AF001504	partial cds	0.4	
~~~~~~~~	AB002384	Human mRNA for KIAA0386 gene, complete cds	0.4	
911	U40145	Mus musculus Mdm2 (Mdm2) gene, complete cds.	0.4	
		Homo sapiens protein phosphatase 1, regulatory subunit 10		
912	NM_002714.1	(PPP1R10) mRNA > :: emb Y13247 HSFB19 Homo sapiens	0.4	
		Mokola virus genes encoding nucleoprotein,	***************************************	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Y09762	phosphoprotein, matrice protein, glycoprotein and	0.4	
	U75187	Arabidopsis thaliana germin-like protein (GLP1) mRNA,	0.4	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z82448	R.prowazekii genomic DNA fragment (clone A503F)	0.39	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z50864	L.delbrueckii sp. lactis plasmid pWS58 DNA, complete	0.39	
917	K03196	Human interferon-beta-3 gene.	0.39	
		Marinococcus halophilus L-2,4-diaminobutyric acid acetyl		
		transferase (ectA) gene, L-2,4-diaminobutyric acid		
		transaminase (ectB) gene, ectoine synthase (ectC) gene,		
918	U66614	putative transposase orfA gene, complete cds, and putative	0.39	
919	L48522	Homo sapiens tuberin (TSC2) gene, exons 6, 7, and 8	0.39	
1000		Caenorhabditis elegans cosmid F23C11, complete sequence	######################################	
www.comencer.com	Z68133	[Caenorhabditis elegans]	0.39	
	U60973		0.39	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	L31398		0.39	
923	X15966	Rabbit DNA for L1Oc4 repeat	0.39	
Ban-((()))-/(()		Caenorhabditis elegans cosmid VZC374L, complete	990-100 (100 (100 (100 (100 (100 (100 (100	
	Z95122	sequence [Caenorhabditis elegans]	0.39	
925	X55036		0.39	
926	U35640		0.39	
	TO THE COLOR OF THE PROPERTY O	Fibroin gene intervening sequence from Bombyx mori (silk		
927	V00094	· · · · · · · · · · · · · · · · · · ·	0.39	
		Streptococcus pneumoniae immunoglobulin A1 protease	<u> </u>	
928	U47687		0.39	
		Caenorhabditis elegans cosmid F28G4, complete sequence		
929	Z93381		0.39	
		Sambucus nigra ribosome inactivating protein precursor	······································	
930	U76524		0.39	
······································	X71600	(*************************************	0.39	

EQ ID	ACCESSION	DESCRIPTION	P VALUI
***************************************		Homo sapiens prostaglandin E receptor 4 sapiens	1 1110
		prostaglandin E2 receptor EP2 subtype mRNA, complete	2000
932	NM 000958.1	cds. > :: gb I36298 I36298 Sequence 1 from patent US	0.39
	Y07648	A.thaliana nit2 gene, nit1 gene and nit3 gene	0.39
	Y13852	Drosophila diplacantha clone D3 inactive Bari-1 family	0.39
	M57851	Human S protein-alpha (PS-alpha) gene, exon 13.	0.39
		Maize chloroplast 3'part of rpoC2 gene, rps2 gene, atpI gene	10.39
936	X52270	and 5'part of atpH gene	0.39
·		Rattus norvegicus taste bud receptor protein TB 567 (TB	0.39
937	U50948	567) gene, complete cds	0.39
		H.sapiens gene PACAP for pituitary adenylate cyclase	10.39
938	X60435	activating polypeptide	0.20
unimener un concenço sence nome.	U12972		0.39
~~~~~	X54709	Tetrahymena thermophila CU428.1VII micronuclear R	0.39
940	A34709	Kluyveromyces lactis BiP gene for BiP/GRP78	0.39
0.41	772260	Human DNA sequence from cosmid 92M18, BRCA2 gene	
941	Z73360	region chromosome 13q12-13	0.39
0.40	T 40044	Plasmid pX01 (from Bacillus anthracis UM23-1) trans-	
	L13841	acting positive regulator (Atx A) gene, complete cds.	0.39
	Z72554	S.cerevisiae chromosome VII reading frame ORF YGL032c	0.39
this is the first own or come on a graph	L24905	Bombyx mori nuclear polyhedrosis virus hr5 gene.	0.39
*******	Z11486	Pinus strobus L. mRNA for pine globulin-1	0.39
~~~~	X70675	M.musculus gene for MERANTES protein	0.39
***************************************	AF010537	Plasmodium falciparum microsatellite TA3 sequence	0.39
948	X78219	D.melanogaster (Barton) SED5 mRNA	0.39
949	D42055	Human mRNA for KIAA0093 gene, partial cds	0.39
	Z30698	Rinderpest virus (RBOK) RNA for RNA polymerase (L)	0.39
951	M64715	Plasmodium falciparum DNA polymerase-delta gene,	0.39
952	Z32774	H.sapiens gene for N-methyl-D-aspartate receptor R1 exons	0.39
		Phanerochaete chrysosporium manganese peroxidase	
953	U70998	isozyme 3 (mnp3) gene, complete cds	0.39
954	Z72884	S.cerevisiae chromosome VII reading frame ORF YGR099w	0.39
······································	S78378	Oct-4 (t12 haplotype) [mice, Genomic, 1191 nt, segment 3	0.39
***************************************	000000000 0000000000000000000000000000	Pyrococcus furiosus ribonucleotide reductase (rnr) gene,	0.37
956	U78098	complete cds	0.38
***************************************		Rattus norvegicus protein serine/threonine kinase CPG16	0.30
957	U78857	*	0.20
	Z54096		0.38
· · · · · · · · · · · · · · · · · · ·	Z72778	S. cerevisiae chromosome VII reading frame ORF YGL256w	0.38
orania and an anti-	D14718		************************************
	D14/10		0.38
061	U09138	Mus musculus peroxisome proliferator activated protein-	
~~~~	**************************************		0.38
······································	U32818		0.38
	X89398	{	0.38
NAME AND	X69440	N.tabacum NTP303 gene	0.38
·····	U62660	Toxoplasma gondii micronemal protein MIC2, complete cds	0.38
······	AE000042	Mycoplasma pneumoniae section 42 of 63 of the complete	0.38
·····	V00190		0.38
·		H.sapiens type X collagen gene	0.38
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U54776		0.38
970 2	Z72888	S.cerevisiae chromosome VII reading frame ORF YGR103w	
		Plasmodium falciparum HMG-like protein Pf16 (Pf16)	
971 t	U97128	7371	0.38
		Human tissue factor gene, complete cds.	

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
*************	J00803	Sheep corticotropin-releasing factor (CRF) mRNA, complet	
*************************	X04310	Rat thymocyte mRNA for 37K chain of CD8 antigen	0.38
<b>***</b>	Y13129	Rickettsia akari fmt gene, partial	0.38
***************************************	X95099	C.albicans mRNA for neutral trehalase	0.38
*********	D83948	Rat adult liver mRNA for S1-1 protein, complete cds	0.38
***************************************		Human DNA sequence from cosmid GG4 from a contig	0.38
978	Z84722	from the tip of the short arm of chromosome 16, spanning	0.38
***************************************	U40604	Listeria monocytogenes ClpC ATPase (mec) gene, complete	0.38
***************************************		Caenorhabditis elegans cosmid M04C3, complete sequence	0.56
980	Z92808	[Caenorhabditis elegans]	0.38
981	X91882	Z.mays ZEMa gene	0.38
982	M26394	M.sexta larval cuticle protein (LCP-14) gene, exon 1.	0.38
***************************************	antinoning in the term of the second of the	Gossypium tomentosum 5S ribosomal RNA gene and non-	0.36
983	U32021	transcribed spacer, clone 11	0.38
*	***************************************	D. melanogaster mRNA coding for a 205K microtubule-	0.30
984	X54061	associated protein (MAP)	0.38
compression and an accompression of the	X14612	Chicken c-myb proto-oncogene 5'region exons 1-5	0.38
***************************************		Sus scrofa immunoglobulin alpha heavy chain constant	0.36
986	U12594	region (IgA C alpha) mRNA, partial cds.	0.38
987	U64841	Caenorhabditis elegans cosmid ZC142.	0.38
988	M93148	Mouse homeobox protein (Hox-1.11) gene, complete cds.	0.38
989	X14049	Canis familiaris RDC4 mRNA for G protein-coupled	0.38
990	X94616	M.musculus mRNA for glycogen synthase	0.38
991	S78378	Oct-4 (t12 haplotype) [mice, Genomic, 1191 nt, segment 3	0.38
	D50091	Drosophila ezoana G-3-P dehydrogenase	0.37
	Y11874	M.musculus uPA gene, promoter sequence	0.37
994	X63628	S.pombe MFm2 gene	0.37
***************************************		Human DNA sequence from cosmid L75B9, Huntington's	
995	Z69652	Disease Region, chromosome 4p16.3	0.37
996	Y08305	L.esculentum lap17.1a gene, promoter region and CDS	0.37
		Human ELK1 pseudogene (ELK2) and immunoglobulin	
	J64453		0.37
	X82286	H.sapiens Fas, Apo-1 gene (exon IX)	0.37
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z48231	E.coli IncQ plasmid pIE639 sat3 gene and aphA gene	0.37
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	L36679	Amycolatopsis methanolica plasmid pMEA300 stf gene,	0.37
1001	X95275	P.falciparum complete gene map of plastid-like DNA	0.37
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	0 00 00 00 00 00 00 00 00 00 00 00 00 0
ana an	Z58565	44a5, forward read cpg44a5.ft1a	0.37
	.14625	Arcobacter skirrowi 16S ribosomal RNA.	0.37
a aran mananana afina	J72236	Dictyostelium discoideum ModA (modA) gene, complete	0.37
1005 3	ζ73940	B.taurus microsatellite DNA INRA190	0.37
	***************************************	Mesocricetus auratus acidic fibroblast growth factor gene, 5'	***************************************
·····	.06092	flank and non-coding exon 1.	0.37
introverse and a second and a second	ζ98918	Human rotavirus gene encoding outer capsid glycoprotein	0.37
	(14448 )	Human GLA gene for alpha-D-galactosidase A (EC	0.37
	AF003533	Homo sapiens cytosolic phagocyte oxidase protein	0.37
1010 X	(67711	O.sativa hsp70 gene for heat shock protein 70	0.37
1011 Z	ann ar an	B.occidentalis gene for 18S rRNA	0.37
1012 Z	12652	R.norvegicus upstream of gene for catechol	0.37
annonenee.	j.	Human papillomavirus type 68, isolate IS362, major capsid	***************************************
1013 U	[45934	protein L1 (L1) gene, partial cds	0.37
1014 U	17084	Human neurofibromin (NF1) gene, promoter region and	0.37
	95668		0.37

Table 2A Nearest Neighbor	(BlastN vs. GenBank)
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EQ ID	4	DESCRIPTION	P VALUI
	X58541	Mink mRNA for plasminogen activator inhibitor type 1	0.37
***********	U48706	Dictyostelium discoideum small aggregate formation	0.37
~~~	M77176	H.sapiens carbonic anhydrase II (CAII) gene, exons 1 and 2.	0.37
***********	Z72888	S.cerevisiae chromosome VII reading frame ORF YGR103v	v 0.37
1020	Y13234	Chironomus tentans mRNA for chitinase, 1695 bp	0.37
	***************************************	Lycopersicon pennellii 2-isopropylmalate synthase (lp-	
*****************	AF004166	ipmsb) mRNA, complete cds	0.37
1022	D85240	Aspergillus niger DNA for isopullulanase precursor,	0.37
	111111111111111111111111111111111111111	Rabbit mRNA for gamma-subunit of phosphorylase kinase	
1023	Y00684	(EC 2.7.1.38)	0.37
		D.discoideum mRNA for 24 kDa protein, homologous to C-	***************************************
1024	X54016	terminal repeat sequence of rhodopsin and synaptophysin	0.37
1025	M22015	Influenza virus type C (C/JJ/50) nonstructural	0.37
1026	M62798	F.ferrugineum 16S ribosomal RNA.	0.37
1027	X76652	M.musculus mRNA for 3f8	0.37
1028	X56047	P. chrysosporium trpC gene for trifunctional polypeptide	0.37
1029	Z74896	S.cerevisiae chromosome XV reading frame ORF YOL154w	0.37
		Mus musculus GTPase-activating protein GAPIII mRNA,	<u> </u>
1030	U20238	complete cds.	0.37
1031	Z54240	L.plantarum pyrimidine biosynthetic operon genes	0.36
	TO CONTRACTOR AND CONTRACTOR OF THE STATE OF	Nicotiana plumbaginifolia beta-(1,3)-glucanase gene for a	0.50
1032	X54742	vacuolar isoform	0.36
		Arabidopsis thaliana 14-3-3-like protein GF14 upsilon	10.30
1033	AF001415	(GRF5) gene, complete cds	0.36
		S.beecheyi genomic DNA with direct repeats and integrated	10.30
1034	X77801	Hepatitis B virus	0.36
	Z84431	D.ligulata internal transcribed spacer 2 DNA	0.36
www.com	X60753	P.sylvestris BBS gene for bibenzyl synthase	0.36
	THE CONTRACT OF SOUTH CONTRACT OF THE CONTRACT	Clostridium josui hemA, hemC, hemD and hemB genes of	0.30
1037	D28503	porphyrin biosynthesis	0.26
	X66131	R. americana mRNA for elongation factor 1-alpha	0.36
	X69601	B.burgdorferi p93 gene (partial)	0.36
	and the second s	Mytilus edulis mitochondrial cytochrome oxidase subunit II	0.36
1040	M83758	(COII) gene, 3' end and NADH dehydrogenase subunit 1	0.06
·····	Z27392	G.raimondii (D61) copia-like reverse transcriptase	0.36
1011		Infectious laryngotracheitis virus US10, US2, protein kinase,	0.36
Service.		UL47, glycoprotein G, ORF5, glycoprotein D, glycoprotein	
1042	U28832	L glycoprotein C. ORFO comes assurated at	0.04
or and the second	AB002384		0.36
	M22345	Human mRNA for KIAA0386 gene, complete cds	0.36
marini marini marini di co	V122343 U16850	Mouse endogenous provirus gag, pol, and env region DNA.	0.36
		Human calmodulin-I (CALM1) mRNA, 3'UTR, partial	0.36
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		a	0.36
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~			0.36
TOTAL PROPERTY OF THE PARTY OF	Z74369		0.36
	S78378		0.36
1020 7	X60686	B.taurus mRNA for transglutaminase	0.36
***	9	Homo sapiens armadillo repeat gene deleted in velo-cardio-	
1051		facial syndrome (ARVCF) mRNA > ::	
1051	√M_001670.1	gb U51269 HSU51269 Human armadillo repeat protein	0.35
		Y.enterocolitica ampC and ampR genes for beta-lactamase	
		and AmpR regulatory protein	0.35
	26270	TT4	0.35

SEQ ID	ACCESSION	DESCRIPTION	P VALU
	***************************************	Listeria monocytogenes internalin (inlC2), inlD, and inlE	
1054	U77368	genes, complete cds	0.35
		Homo sapiens intergenic locus pYNZ32 variable number	
1055	M62750	tandem repeat (VNTR) sequence associated with Huntington	0.35
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		Slime mold (D.discoideum) mRNA complementary to the	
1056	K03073	right inverted terminal repeat of DIRS-1, clone pLZ12.	0.35
1057	U78025	Sus scrofa microsatellite S0399 sequence	0.35
······	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Drosophila melanogaster transformer (tra) gene, complete	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
1058	M17478	cds, and non-sex-specific transformer processed pseudogene	0.35
		Homo sapiens complement component 2 (C2) gene allele b,	
1059	L09708	exons 10 through 18 and complete cds	0.35
~~~~	Z16708	H. sapiens (D6S275) DNA segment containing	0.35
tree researches and the annual companions	M74201	Rabbit cytochrome P450IIC4 (CYP2C4) gene, exons 2 and	0.35
*****************************	,	Mesembryanthemum crystallinum inositol methyltransferase	· · · · · · · · · · · · · · · · · · ·
1062	U63634	(Imt1) gene, complete cds	0.35
~~~~~		Oryctolagus cuniculus calcitonin receptor isoform mRNA,	ļ
1063	U73126	complete cds	0.35
	X52614	Maize chloroplast rps15 gene for ribosomal protein S15	0.35
····	X52614	Maize chloroplast rps15 gene for ribosomal protein S15	0.35
	X57520	L.micdadei DNA for gro ELS operon	0.35
***************************************	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	Pyrodictium occultum polB gene for DNA polymerase II	**************************************
1067	D38574	(family B) (alpha-like DNA polymerase), complete cds	0.35
		Caenorhabditis elegans cosmid E02H1, complete sequence	10.33
1068	Z47075	[Caenorhabditis elegans]	0.35
annonano angrangangangan da di	Y10686	C.capreolus DNA, tandem repeat region	0.35
	***************************************	Blaberus giganteus mitochondrion 16S ribosomal RNA,	10.55
1070	U <b>17</b> 771	partial sequence	0.35
		Oryctolagus cuniculus sodium/dicarboxylate cotransporter	
1071	U12186	mRNA, partial cds.	0.35
~~~~~	L25677	Schizosaccharomyces pombe cdc42+ (Cdc42p) gene,	0.35
1073	X63628	S.pombe MFm2 gene	0.35
~~~~~	D67071	Rat gene for regucalcin, exon1 (non-coding exon)	0.35
***************************************	X93218	Impatiens necrotic spot virus L gene	0.35
	U16301		0.35
	M37277	Human Ig germline H-chain D-region genes, partial cds.	0.35
	X14448	Human GLA gene for alpha-D-galactosidase A (EC	0.34
mannonnonnonnonnonnonnon	X79930	O.gibsoni Tbb gene	0.34
·····		H.sapiens rearranged immunoglobulin lambda light chain V-	
1080	Z37352	region (VI-3)	0.34
<b>wasanananananananananana</b>	X59013		0.34
1082	X90761	Homo sapiens hHa2 gene	0.33
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	L23498	Bovine microsatellite repeats	0.33
1084	Z18361	O.aries trichohyalin	0.33
·····	L18785		0.33
erennen er	X96685	B.burgdorferi cell division genes	0.33
	M58509		0.33
····	X70809	C.reptans gene for 18S ribosomal RNA	0.32
www.commons.com	D38116	·	0.32
		Human immunodeficiency virus type 1 isolate 583-78	~- <b>~-</b>
1090	U00310	·	0.31
	······	Homo sapiens (subclone H8 10 g5 from P1 35 H5 C8) DNA	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	**************************************	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	0.3

SEQ ID	ACCESSION	DESCRIPTION	P VALU
	M80215	Streptococcus pneumoniae uvs402 protein gene, complete	0.22
***	M98776	Human keratin 1 gene, complete cds	0.22
****	X66313	H.sapiens GLUDP2 gene (exon 2)	
	AB001025	Homo sapiens mRNA for brain ryanodine receptor, complete	0.21
	AE000956	Archaeoglobus fulgidus section 151 of 172 of the complete	торитов точно гинического основных
	Z46268	Simian herpesvirus B DNA for glycoprotein G	0.21
************************	X99403.1	N.tabacum mRNA for defensin	0.21
1100	X77403.1	Drosophila pseudoobscura EP arrangement, BC p430#4	0.21
			**
		pAP5 clone alpha-amylase (amy1) gene, complete cds. > ::	***
1101	U20335	gb U20337 DPU20337 Drosophila pseudoobscura HI	
	X04077	arrangement, Mex a-176x pAP9 clone alpha-amylase (amy1)	
PETERSON DECIMAL PERSON NAMED AND ADDRESS OF THE	TOP ACCOUNTS AND AND AN ADDRESS OF PROPERTY OF PROPERTY OF THE PARTY O	Potato patatin pseudogene (SB6B)	0.2
1103	L11236	Rattus rattus mRNA sequence.	0.2
1104	37.40.5.40	H.sapiens mRNA for tre oncogene (clone 213) > ::	
1104	X63547	gb 176205 176205 Sequence 9 from patent US 5691147	0.2
440-		Human liver glucokinase (ATP:D-hexose 6-	
1105	M69051	phosphotransferase) mRNA, complete cds.	0.2
		Rattus norvegicus synapse-associated protein 97 mRNA,	
1106	U14950	complete cds.	0.2
		Homo sapiens PRKC, apoptosis, WT1, regulator apoptosis	
	NM_002583.1	response protein par-4 mRNA, complete cds	0.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X66313	H.sapiens GLUDP2 gene (exon 2)	0.2
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U48485	Human skeletal muscle ryanodine receptor gene	0.2
gamana a manana a m	AF018116	Dendroica pityophila cytochrome oxidase I protein, partial	0.2
1111	X87108	S.cerevisiae CKI3 gene	0.19
VOI.		E.coli retron Ec67 DNA encoding reverse transcriptase and	<u> </u>
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M55249	Dam methylase functions.	0.19
manar manar manar di sa	L06898	Actinomyces viscosus sialidase (nanH) gene, complete cds.	0.19
·	Y13631	Clostridium botulinum P-21, P-47 ntnh, bonT genes	0.18
talaanin aan oo	L24904	Bombyx mori nuclear polyhedrosis virus hr4R gene.	0.18
***************************************	M85049	Drosophila melanogaster brahma protein mRNA, complete	0.18
1117	X60777	Shigella dysenteriae ipaBCD genes	0.17
		Pseudoxycheila bipustulata mitochondrial large subunit 16S	18 10 10 10 10 10 10 10 10 10 10 10 10 10
1118	L42936	ribosomal RNA (16S rRNA) gene	0.17
1119	AE001359	Chlamydia trachomatis section 86 of 87 of the complete	0.17
		yeast(s.cerevisiae) histone 3 gene (h3) fused with e.coli lacz	
1120	X00899		0.17
and the same of th		Staphylococcus aureus prolipoprotein signal peptidase (lsp)	
1121		,	0.17
1122 2	Z28050	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	0.16
	THE PARTY OF THE P	Mus musculus fragile X mental retardation syndrome protein	·····
1123 1	L23971	I	0.16
1124 2	<b>Z</b> 98560	and the second s	0.15
	J89258		0.15
·····································	₹62522	·	0.15
	₹74504		0.15
····	·····	^	0.15
	~~~		
<del>marana makanga kangina</del>			0.15
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Blepharoneura manchesteri 16S ribosomal RNA gene,	0.15
1131 T			0.15
*****	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Leptocarabus procerulus mitochondrial DNA for NADH	0.15

SEQ ID	ACCESSION	DESCRIPTION	P VALU
1133	L35272	Glycine max heat shock protein (SB100) mRNA, complete	0.15
		Legionella pneumophila rpoD operon LporfX, LpdnaG, and	
1134	U63641	LprpoD genes, complete cds	0.15
1135	X72186	H.sapiens C6 gene, exons 10, 11 and 12	0.15
1136	D45893	Neurospora crassa acr-2 DNA, complete cds	0.15
1137	L47106	Neurospora crassa kinesin (NKIN) mRNA, complete cds	0.15
	······································	Phytophthora megasperma mitochondrial ORF152, complete	
		cds, cytochrome c oxidase subunit I (cox1) gene, complete	***
1138	L04457	cds, cytochrome c oxidase subunit II	0.15
		Phytophthora megasperma mitochondrial ORF152, complete	
		cds, cytochrome c oxidase subunit I (cox1) gene, complete	
1139	L04457	cds, cytochrome c oxidase subunit II	0.15
1140	U12539	Schizosaccharomyces pombe scd2 (scd2) gene, complete	0.15
1141	U12539	Schizosaccharomyces pombe scd2 (scd2) gene, complete	0.15
	AB001901	Homo sapiens PACE4 gene, exon 4-7	0.15
		Maize sucrose phosphate synthase mRNA, complete cds. > ::	
1143	M97550	emb A17878.1 A17878 Synthetic SPS sequence >	0.15
	AB001901	Homo sapiens PACE4 gene, exon 4-7	0.15
	X07438	Human DNA for cellular retinol binding protein	0.14
	1207.100	H.sapiens CpG island DNA genomic Mse1 fragment, clone	<u> </u>
1146	Z57339	171g3, reverse read cpg171g3.rt1a	0.14
***************************************	X05034	Rat C2A gene for prostatic binding protein (PBP)	0.14
u reconstruction and a second	U55712		0.14
***************************************	M20273	<u> </u>	0.14
and the second second second second	J01764	<u>,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,</u>	0.14
	X61109	<u>, and an announce and an announce and an announce and an announce and an an announce and an announce and an an</u>	0.14
1101		S.hominis (972) genes for potential ABC transporter and	ļ
1152	Z30587	potential membrane spanning protein	0.14
		Babanki virus polymerase nsP4 gene, partial cds virus	
1153	U60400	polymerase nsP4 gene, partial cds	0.14
	U67478	Methanococcus jannaschii section 20 of 150 of the complete	
	Z71641	S.cerevisiae chromosome XIV reading frame ORF	0.14
www.co.co.co.co.co.co.co.co.co.co.co.co.co.	X53522	M.musculus VL1 gene (1G-5)	0.14
	Z21753	O.tanganicae mitochondrion genes for tRNA-Thr	0.14
1137	221733	Carthamus tinctorius glycerol-3-phosphate acyltransferase	0.11
1158	L33841	,	0.14
1130		Glossiphonia complanata cytochrome c oxidase subunit I	
1150	AF003277	(COI) gene, partial cds	0.14
1137	A1 003211	Oxytricha fallax transposon TBE1, insertion fal6, 42 kDa	U.1 T
1160	U89035	transposase gene, partial cds	0.14
1100	009033	Streptococcus pneumoniae beta-N-acetylhexosaminidase	U. I T
1161	L36923	(strH) gene, complete cds	0.14
1101		Dictyostelium discoideum AX2 protein tyrosine kinase gene,	
1162	U64830	complete cds.	0.14
1102	201030	Zea mays T cytoplasm male sterility restorer factor 2 (rf2)	
1162	U43082	mRNA, complete cds	0.14
	Z26492	T.repens TrMT1A mRNA for metallothionein-like protein	0.14
	X83683	V.sativa mRNA for early nodulin 40	0.14
war incompanies and the contract of the contra	THE PROPERTY AND ADMINISTRATION OF THE PROPERTY AND ADMINISTRATION OF THE PARTY AND ADMINISTRA	Homo sapiens DNA for cyclophilin 40, complete cds	0.14
1100	D63861	Japanese jack bean clone CgHMGY1 DNA for high mobility	
1167	D86594	group protein, complete cds	0.14
110/	し 00374	group brotem, comblete cas	V.14

Table 2A Nearest	Neighbor	(BlastN vs.	GenBank)
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SEQ ID	ACCESSION	DESCRIPTION	P VALU
44.0		Trypanosoma rangeli kinetoplast DNA sequence with	
	L28038	conserved sequence blocks	0.14
1170	U72057	Mus musculus chloride channel regulator (Icln) gene, exon 1	0.14
		Mesocricetus auratus isopentenyl diphosphate:dimethylallyl	resolution a
	AF003836	diphosphate isomerase mRNA, complete cds	0.14
	X78422	A.thaliana HYP1 mRNA	0.14
**********************	U28042	Human DEAD box RNA helicase-like protein mRNA,	0.14
***************	U01070	Mycoplasma flocculare Ms42 5S rRNA gene.	0.14
~~~~~	U54776	Human NTT gene, L1, Alu, and MER 38 repeat regions	0.14
**********************	U40375	Mus musculus Supt6h mRNA, complete cds	0.14
	X71083	G.max gene for coproporphyrinogen oxidase	0.14
1178	D86253	Human MHC (HLA) DRB intron 1 DNA, partial sequence	0.14
		Synechococcus sp. 6-phosphogluconate dehydrogenase	
	M55002	gene, complete cds.	0.14
1180	X14036	Tomato cab-7 gene for type II chlorophyll a/b-binding	0.14
1181	L35272	Glycine max heat shock protein (SB100) mRNA, complete	0.14
1182	D50931	Human mRNA for KIAA0141 gene, complete cds	0.14
1183	L35272	Glycine max heat shock protein (SB100) mRNA, complete	0.14
1184	X05034	Rat C2A gene for prostatic binding protein (PBP)	0.14
1185	X89553	Echovirus 26 DNA for VP2 capsid protein gene	0.14
		Caenorhabditis elegans cosmid F07B10, complete sequence	\$
1186	Z77656	[Caenorhabditis elegans]	0.14
1187	Z84739	H.sapiens flow-sorted chromosome 6 HindIII fragment,	0.14
		Homo sapiens retinaldehyde-binding protein (CRALBP)	<b>~~~~~</b>
1188	L34219	gene, complete cds.	0.14
1189	M30168	D.melanogaster nested repetitive sequences F and G,	0.14
		Perissodus microlepis T32a mitochondrion NADH	
1190	U07260	dehydrogenase subunit 2 gene, complete cds	0.14
1191	X15308	H.sapiens NF-H gene, exon 3	0.14
1192	J02763	Human calcyclin gene, complete cds.	0.14
1193	M76175	Mouse elongation factor 2 pseudogene, complete cds.	0.14
1194	X67735	M.musculus mas gene for Mas proto-oncogene	0.14
	f. A. A. Territorio, Ser	Homo sapiens proteasome (prosome, macropain) 26S	(1970-1984 <del>)</del> 1980-1980-1980-1980-1980-1980-1980-1980-
		subunit, non-ATPase, 12 (PSMD12) mRNA > ::	
1195	NM 002816.1	dbj AB003103 AB003103 Homo sapiens mRNA for 26S	0.14
1196	U20754	Felis catus chromosome D2 mtDNA tandem repeat Numt	0.14
		Human myotonin protein kinase (Mt-PK) gene, complete	teristi assentare service economico es-
1197	L00727		0.14
1198	M57750	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	0.14
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1199	X03000	Adenovirus type 7 genome left end (0.0 to 31 %)	0.14
1199	X03000		0.14
1199	X03000	Homo sapiens regulator of nonsense transcripts 1 (RENT1)	0.14
	dalaharan sain sa mengan mengel kerencerangan pangan pangan mengan menjada menjada menjada menjada menjada men	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of	TO THE PERSON AND ADDRESS OF THE PERSON ADDRESS OF THE PERSON AND ADDRESS OF THE PERSON AD
1200	NM_002911.1	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds	0.14
1200 1201	NM_002911.1 Z71367	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S.cerevisiae chromosome XIV reading frame ORF	0.14 0.14
1200 1201 1202	NM_002911.1 Z71367 J04809	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S.cerevisiae chromosome XIV reading frame ORF Human cytosolic adenylate kinase (AK1) gene, complete	0.14 0.14 0.14
1200 1201 1202 1203	NM_002911.1 Z71367 J04809 X75014	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S.cerevisiae chromosome XIV reading frame ORF Human cytosolic adenylate kinase (AK1) gene, complete M.musculus Phox2 mRNA for homeodomain protein	0.14 0.14 0.14 0.14
1200 1201 1202 1203 1204	NM_002911.1 Z71367 J04809 X75014 Y12259	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S.cerevisiae chromosome XIV reading frame ORF Human cytosolic adenylate kinase (AK1) gene, complete M.musculus Phox2 mRNA for homeodomain protein R.norvegicus mRNA for Kir3.1 protein	0.14 0.14 0.14 0.14 0.14
1200 1201 1202 1203 1204 1205	NM_002911.1 Z71367 J04809 X75014 Y12259 X81981	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S.cerevisiae chromosome XIV reading frame ORF Human cytosolic adenylate kinase (AK1) gene, complete M.musculus Phox2 mRNA for homeodomain protein R.norvegicus mRNA for Kir3.1 protein L.helveticus plasmid pLH2 DNA (4 ORFs)	0.14 0.14 0.14 0.14 0.14 0.14
1200 1201 1202 1203 1204 1205 1206	NM_002911.1 Z71367 J04809 X75014 Y12259 X81981 AB001901	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S. cerevisiae chromosome XIV reading frame ORF Human cytosolic adenylate kinase (AK1) gene, complete M.musculus Phox2 mRNA for homeodomain protein R.norvegicus mRNA for Kir3.1 protein L.helveticus plasmid pLH2 DNA (4 ORFs) Homo sapiens PACE4 gene, exon 4-7	0.14 0.14 0.14 0.14 0.14 0.14 0.14
1200 1201 1202 1203 1204 1205 1206	NM_002911.1 Z71367 J04809 X75014 Y12259 X81981	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S.cerevisiae chromosome XIV reading frame ORF Human cytosolic adenylate kinase (AK1) gene, complete M.musculus Phox2 mRNA for homeodomain protein R.norvegicus mRNA for Kir3.1 protein L.helveticus plasmid pLH2 DNA (4 ORFs) Homo sapiens PACE4 gene, exon 4-7 Human mRNA for KIAA0076 gene, complete cds	0.14 0.14 0.14 0.14 0.14 0.14
1200 1201 1202 1203 1204 1205 1206	NM_002911.1 Z71367 J04809 X75014 Y12259 X81981 AB001901	Homo sapiens regulator of nonsense transcripts 1 (RENT1) mRNA > :: gb U65533 HSU65533 Human regulator of nonsense transcript stability (RENT1) mRNA, complete cds S. cerevisiae chromosome XIV reading frame ORF Human cytosolic adenylate kinase (AK1) gene, complete M.musculus Phox2 mRNA for homeodomain protein R.norvegicus mRNA for Kir3.1 protein L.helveticus plasmid pLH2 DNA (4 ORFs) Homo sapiens PACE4 gene, exon 4-7	0.14 0.14 0.14 0.14 0.14 0.14 0.14

Table 2A Nearest Neighbor (BlastN vs. GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
		Caenorhabditis elegans cosmid C44E1, complete sequence	
	Z92779	[Caenorhabditis elegans]	0.13
1210	X14384		0.13
		Bacillus thuringiensis cryIC-related gene sequence. > ::	
		gb I82521 I82521 Sequence 1 from patent US 5712248 > ::	
	M97880	gb I93759 I93759 Sequence 1 from patent US 5731194	0.13
1212	X58049	H.sapiens DNA for ferredoxin repeat region	0.13
		Vibrio cholerae ADP-L-glycero-D-mannoheptose-6-	
1213	U47542	epimerase (rfaD) gene, complete cds.	0.13
		Sheep (clone C5-2) MHC class II cell surface glycoprotein	
		OLA-DRB (OLA-DRB) pseudogene, exon 4. > ::	
1214	M90758	gb S83918 S83914S3 DRB=MHC class II B {pseudogene}	0.13
1215	U67478	Methanococcus jannaschii section 20 of 150 of the complete	0.13
1216	U90889	Mus musculus transketolase (TKT) gene, partial cds	0.13
		C.elegans heat shock protein genes (hsp16-48 and hsp16-1),	
1217	K03273	complete cds.	0.13
1218	D90736	Escherichia coli genomic DNA. (22.6 - 23.0 min)	0.13
1219	X63546	H.sapiens mRNA for tre oncogene (clone 210)	0.13
		Anas platyrhynchos mitochondrial complete transfer RNA-	\$
		Glu, transfer RNA-Phe, transfer RNA-Val, transfer RNA-	
1220	L16770	Leu, 12S ribosomal RNA, and 16S ribosomal RNA genes	0.13
	U <b>9441</b> 0	Dictyostelium discoideum plasmid Ddp6 Rep protein	0.13
1222	AC001083	Homo sapiens (subclone 2 a6 from BAC H75) DNA	0.13
		Caenorhabditis elegans cosmid F57A10, complete sequence	
1223	Z96048	[Caenorhabditis elegans]	0.13
~~~~~~~~	M92914	Drosophila virilis mastermind gene, complete cds	0.13
	Z82961	Bacterial sp. partial 16S rRNA gene (clone group T4A)	0.13
1226	U66535	Human beta4-integrin (ITGB4) gene, exons	0.13
. B. D. C.	<del>in man en men en e</del>	Homo sapiens epiregulin (EREG) mRNA > ::	\$0.000 mm.
1227	NM 001432.1	dbj D30783 D30783 Homo sapiens mRNA for epiregulin,	0.13
		P.tetraurelia gamma1-51D immobilisation antigen gene, 3'	
1228	X96468	coding and non-coding region	0.13
e paragonales apares a representado de la constancia de la constancia de la constancia de la constancia de la c	X05918	Kluyveromyces fragilis beta-glucosidase gene	0.13
	Z48243	A.thaliana PARP mRNA for PARP protein	0.13
	X56276	Human Hut 2 End gene	0.13
	X55318	Mus musculus Hox-3.2 gene	0.13
		Phalacrocorax pelagicus cytochrome b gene, mitochondrial	
1233	U90009	gene encoding mitochondrial protein, partial cds	0.13
	L76571	Homo sapiens nuclear hormone receptor (shp) gene, 3' end	0.13
		Human c-jun gene, promoter region with flanking	
1235	U60581	evolutionary conserved sequences	0.13
	Z26886	B.mori gene for Nd-sD mutant fibroin light chain	0.13
		Canis familiaris Kv1.2 delayed rectifier K+ channel mRNA,	
1237	L19740	complete cds	0.13
1257		Glycine max ribosomal protein S16 (rps16) gene, partial cds,	
		beta-carboxyltransferase (accD), photosystem I component	
		(psaI), ORF 202 protein (ORF 203), ORF 151 protein (ORF	
1238	U26948	151), ORF 103 protein (ORF 103), ORF 229 precurso	0.13
1230	U & U / T U	Rattus norvegicus A-kinase anchoring protein AKAP 220	U.1J
1220	1148788	mRNA, complete cds	0.13
and the second s	U48288 X70064	P.deltoides gene for poplar bark storage protein	0.13
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	L22857	Colletotrichum gloeosporioides pectin lyase (pnlA) gene,	0.13
	1.4.40.17	Concloudending stocosportorides decini ivase (dina) gene.	U.13

ACCESSION	DESCRIPTION	P VALU
X82584	M.musculus mRNA for immunoglobulin heavy chain, V	0.13
M76713		0.13
X57075		0.13
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Z79998		0.13
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AF009074		0.13
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Z57339		0.13
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1176524		0.13
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AF000582		0.13
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Z71367	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	0.13
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U38292	Human microtubule-associated protein 1a (MAP1A) mRNA,	0.13
	X82584  M76713 X57075 Z17201 D50931 D80002 M58600  Z79998 X82868  AF009074 M26716  Z57339  U76524  AF000582  D12917 X70326 X82237 U10116 X73293  U01766 X79706 U45957 X66313 X07946 X07870 X15308 Z22551 X89398 D10654 U10516 X57075 U71193 X69908 M81322  X78401  X78401 Y09852	M.musculus mRNA for immunoglobulin heavy chain, V Spodoptera frugiperda 16S rRNA gene, Val-tRNA, and Leu- tRNA genes, and ND-1 protein gene, S' end.  M76713 H.sapiens FGF6 gene Z17201 H. sapiens (DXS1003) DNA segment containing D50931 Human mRNA for KIAA0141 gene, complete cds D80002 Human mRNA for KIAA0180 gene, partial cds M58600 Human mRNA for KIAA0180 gene, partial cds Human DNA sequence from cosmid B5E3 on chromosome Z2 Contains CpG island, EST X82868 G.gallus microsatellite DNA (63G2) Hepatitis C virus genomic RNA, 3' nonstranslated region, partial sequence. clone #19 M26716 Rat cAMP phosphodiesterase mRNA, complete cds., clone H.sapiens CpG island DNA genomic Mse1 fragment, clone Z57339 171g3, reverse read cpg171g3-rt1a Sambucus nigra ribosome inactivating protein precursor mRNA, complete cds Mus musculus nuclear receptor coactivator protein 2 mRNA, complete cds Yeast gene for a component of the pheromone signal transduction pathway, complete cds X70326 H.sapiens MacMarcks mRNA Schinensis RAPD DNA (523 bp) U10116 Human superoxide dismutase (SOD3) gene, complete cds. X73293 M.vannielli genes rpoH, rpoB and rpoA Mycoplasma genitalium major adhesion protein MgPa gene, partial cds X79706 C.aietinum capr1 mRNA for pathogenesis-related protein U45957 Nicotiana alata SA2-RNase precursor gene, complete cds. X76931 X76946 Yeast plasmid DNA coding for RNA polymerase subunit X07870 Drosophila melanogaster bicoid gene bcd X15308 H.sapiens Kip-H gene, exon 3 X222551 H.sapiens FGF6 gene U11193 Mus musculus Rpl23 matrix-attachment region Mus mu

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
1283	Z71367	S.cerevisiae chromosome XIV reading frame ORF	0.13
1284	M87623	Babesia bovis variable antigen 1 mRNA, 5' end.	0.13
1285	M91452	Sus scrofa ryanodine receptor (RYR1) gene, complete cds.	0.13
1286	M91452	Sus scrofa ryanodine receptor (RYR1) gene, complete cds.	0.13
1287	M87623	Babesia bovis variable antigen 1 mRNA, 5' end.	0.13
1288	X96616	P.primaurelia gene encoding 156D surface antigen	0.13
1289	Z33222	M.capricolum DNA for CONTIG MC303	0.12
1290	X52626	Phaseolus vulgaris gene for alpha-phaseolin	0.12
1291	L32699	Paracentrotus lividus glutamine synthetase (GS) mRNA,	0.12
1292	M63669	Dog arginine esterase gene, complete cds.	0.12
1293	X75653	A.longa plastid genes for ribosomal proteins and tRNAs	0.12
1294	Z11839	T.maritima nusG gene and genes for ribosomal proteins	0.12
1295	AB005744	Perilla frutescens DNA for 1-limonene synthase, complete	0.12
1296	X84904	Plasmodium falciparum encoding Pfg27/25	0.12
		Drosophila melanogaster ribosomal protein S6 gene and two	0.12
1297	L02074	potential alternatively spliced proteins, complete cds	0.12
	**************************************	Human DNA sequence from phage pTEL from a contig from	; 0.12 /************************************
www.		the tip of the short arm of chromosome 16, spanning 2Mb of	
1298	Z84812 ·	16p13.3 Contains ESTs	0.12
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X95276	P.falciparum complete gene map of plastid-like DNA	0.12
		Homo sapiens epiregulin (EREG) mRNA > ::	[U.12
1300	NM 001432.1	dbj D30783 D30783 Homo sapiens mRNA for epiregulin,	0.12
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X84726	M.musculus neurocan gene	0.12
······································	Z35810	S.cerevisiae chromosome II reading frame ORF YBL049w	0.12
marana a amanana ang a	L01655	Plasmodium falciparum T9/106 triosephosphate isomerase	0.12
***************************************	U36796		0.12
varaneen va	U22361	Mus musculus presynaptic protein SNAP-25 (Snap-25) gene,	0.12
1303		Saccharomyces cerevisiae Rlr1p (RLR1) gene, complete cds. Plasmodium falciparum (clone Pfg377 [PfsXLX]) DNA	0.12
1306 1	L04161	sequence, complete cds	0.10
manana manana manana kata da k	U32768	Haemophilus influenzae Rd section 83 of 163 of the	0.12
1507			0.12
1308 1	D28808	Mycoplasma capricolum mtlA and gyrB genes for DNA	
13001	D20000	gyrase subunit B and mannitol-specific phosphotransferase	0.12
1300 1	L05920	Human constitutively expressed serum amyloid A protein	
************************	M96642	(SAA4) gene, exons 1 through 4.	0.12
	······································	Paramecium tetraurelia P126 repetitive element.	0.12
1311	J01/24	Homo sapiens (subclone 2_a3 from P1 H69) DNA sequence	0.12
1212	VE010221	Tenebrio molitor thermal hysteresis protein isoform YL-3	
······································	AF010331	(	0.12
1313	J <b>74496</b>	Human chromosome 4q35 subtelomeric sequence	0.12
1214 T	117205	Haemophilus influenzae dppB, dppC, dppD, dppF, isn, artP,	
····	J17295	artI/J, artQ, and artM genes, complete cds, and opa gene,	0.12
	ζ53921	A. thaliana T-DNA insertion genomic target sequence 621-	0.12
1316 2	₹75653	A.longa plastid genes for ribosomal proteins and tRNAs	0.12
1015	T000 4	Human variant urokinase plasminogen activator receptor	
1317 (			0.12
4		hyaluronan-binding protein=hepatocyte growth factor	
1318 S		activator homolog [human, plasma, mRNA, 2408 nt]	0.12
1319 L	J <b>21164</b>	Human dopamine D5 receptor gene, 5' flanking and	0.12
* October 1990	According to	Feline herpesvirus 1 immediate early protein, glycoprotein	**************************************
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	F022391	· · · · · · · · · · · · · · · · · · ·	0.12
1321 N	174569	Clostridium acetobutylicum heat shock protein	0.12
		Drosophila virilis hunchback (hb) gene for zinc-finger	
1322 X		protein transcription factor	

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
	U67478	Methanococcus jannaschii section 20 of 150 of the complete	
	X77052	Entomopoxvirus gene for spherulin	0.12
		Saccharomyces douglasii mitochondrial cytochrome c	0.12
1325	M97514	oxidase subunit I (COXI) gene, complete cds	0.12
	***************************************	H.sapiens DNA for the upstream regulatory region of the c-	0.12 1
1326	X56495	erbB2 gene	0.12
~~**	D61398	a thiga ann a gana ann an an an ann an an an an an an a	0.12
1361	D01376	Caenorhabditis elegans cosmid T22H2, complete sequence	* U. I Z
1328	Z81595	[Caenorhabditis elegans]	0.12
*********************	D61398	Human MSH3 gene, exon3	0.12
1347	D01330	Human DNA sequence from cosmid B5E3 on chromosome	0.12
1220	770000	-	0.12
1330	Z79998	22 Contains CpG island, EST	0.12
1221	NTM 001200 1	Homo sapiens cold inducible RNA-binding protein (CIRBP)	0.13
1331	NM_001280.1		0.12
1222	740077	Caenorhabditis elegans cosmid C33D3, complete sequence	0.13
	Z49867	[Caenorhabditis elegans]	0.12
1333	X14036	Tomato cab-7 gene for type II chlorophyll a/b-binding	0.12
1224	A TO 1 2 0 0 0	Sambucus nigra ribosome inactivating protein precursor	
	AF012899	mRNA, complete cds	0.12
1335	U19159	Human P protein (P) gene, exon 8	0.12
		H.sapiens mRNA for 2'-5' oligoadenylate binding protein >	3
1226	*****	:: gb L24115 HUMAAZ Human DNA fragment. > ::	
1336	X74987	emb A49723.1 A49723 Sequence 2 from Patent	0.12
		Dictyostelium discoideum 34 kDa actin binding protein	
1005	7700110	gene, complete cds > :: emb Z50156 DD30KDABP	
1337	U32112	D.discoideum gene for 34 kD actin binding protein	0.12
1220	1.65000	Synechococcus sp. 6-phosphogluconate dehydrogenase	0.10
	M55002	gene, complete cds.	0.12
1339	AF019225	Homo sapiens apolipoprotein L mRNA, complete cds	0.12
12.40	*********	Mycobacterium smegmatis diptheria toxin repressor	
	U14190	homolog (dtxR) gene, complete cds.	0.12
rano, <del>Nico, enconsociones con con</del> con esciente con con c	J05276	Rat 5-hydroxytryptamine-1a receptor (5-HT-1a) gene,	0.12
	AE000603.1		0.12
~~ <del>~~~~</del>	X15308	H.sapiens NF-H gene, exon 3	0.12
1344	L35600	Homo sapiens DNA sequence.	0.12
		Human DNA sequence from cosmid 92M18, BRCA2 gene	
1345	Z73360	region chromosome 13q12-13	0.12
		Homo sapiens green cone photoreceptor pigment gene, 5'	
1346	U93721	flanking region	0.12
		Bacteriophage P22 right operon, orf 48, replication genes 18	
		and 12, nin region genes, ninG phosphatase, late control	
·····	X78401	gene 23, orf 60, complete cds, late control region, start of	0.12
1348	Z71367	S.cerevisiae chromosome XIV reading frame ORF	0.12
1000		Mus musculus putative transmembrane receptor IL-1Rrp	
200		mRNA, complete cds > :: gb AR016448 AR016448	
and the second second	U43673	Sequence 3 from patent US 5776731	0.12
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X72863	A.thaliana TMKL1 mRNA	0.12
ang merang ang ang ang ang ang ang ang	U13769	Vibrio sp. ppGpp synthetase I (relA) gene, complete cds.	0.12
1352	M32732		0.11
		Xenopus laevis XK81A1 keratin gene > ::	
approximately and the second		emb X04804 XLXK81A1 Xenopus laevis DNA for stage-	
1353	X04668	specific epidermal type I keratin A1 (embryo- and larval-	0.11

SEQ II	ACCESSION	DESCRIPTION	DATATE
	- 11002001011	Human DNA sequence from PAC 293L6 on chromosome	P VALU
135	54 Z82197	22, complete sequence [Homo sapiens]	
		Cucumis sativus ORF 1, chitinase, and ORF 3 genes,	0.11
		complete cds > :: gb I38466 I38466 Sequence 36 from pate	una unana
		119 5614205 > chire could be could be compared to the could be	nt
	***************************************	US 5614395 > :: gb I56941 I56941 Sequence 36 from pater	ıt
135	5 M84214	US 5650505 > :: gb I59807 I59807 Sequence 36 from pater	\$
	6 Y08503	US 5654414 > :: gb I75134 I7513	0.11
POTOTOTO OPENIA LA LA COLONIA MANAGEMENTA	ti este la como de este en	F.domesticus mitochondrial 12S rRNA gene	0.11
	7 L03286	Hamster P-glycoprotein class I (pgp1) gene exons 1-2.	0.11
133	8 X57171	D.caryophyllus CARSR12 gene	0.11
		Mus musculus ATPase mRNA-ampifying genomic DNA	
125		MOR6.5 sequence. > :: gb S55685 S55685	vroon v
135	9 L07307	MOR6.5=ouabain resistance gene {repeat sequence} [mice,	0.11
	»	Caenorhabditis elegans cosmid K01F9, complete sequence	
Market Anna Control Control Control	0 <b>Z22</b> 175	[Caenorhabditis elegans]	0.1
	1 Z11839	T.maritima nusG gene and genes for ribosomal proteins	0.1
	2 L36857	Pisum sativum GTP-binding protein (IAP86) mRNA,	0.096
******************	3 AF000994	Homo sapiens ubiquitous TPR motif, Y isoform	0.07
1364	4 U61950	Caenorhabditis elegans cosmid C45E5	0.069
		Mus musculus putative breast/ovarian cancer susceptibility	
<del>~~~</del>	U32446	protein homolog (Brca1) mRNA, complete cds.	0.068
1366	6 AB001914	Homo sapiens PACE4 gene, exon 23-25, complete cds	0.068
1367	7 L20934	Anopheles gambiae complete mitochondrial genome	0.066
1368	3 AJ001700	Mus musculus mRNA for neuroserpin	0.065
1369	AJ001700	Mus musculus mRNA for neuroserpin	0.065
1370	AF106932	Drosophila melanogaster plexin A (plexA) mRNA, complete	0.003
****************		Populus kitakamiensis cyp73b gene for cinnamic acid 4-	0.003
1371	D82813	hydroxylase, partial cds	0.065
	V01087	Hemagglutinin gene of influenza virus strain	0.065
*****	AE001178	Borrelia burgdorferi (section 64 of 70) of the complete	0.065
************************	X80199	H.sapiens MLN51 mRNA	0.064
	AF042384	Homo sapiens BC-2 protein mRNA, complete cds	0.064
Mechanism consider and again and	AJ223485	Enoploteuthis higginsi mitochondrial 16S rRNA	0.064
****	AF086094	Homo sapiens full length insert cDNA clone YZ87H06	0.063
********************	AF017027	A frican swine fover views leaving have 1 (00P)	0.061
		African swine fever virus lectin homolog (8CR) gene,	0.061
		Bordetella bronchiseptica electron transfer flavoprotein	· ANDROIS
1379	U56084	alpha subunit (etfA) gene, partial cds, and exogenous ferric	anamata.
	M33387	siderophore receptor (bfrA) gene, complete cds	0.06
1300	14133307	Human debrisoquine 4-hydroxylase (CYP2D8P) and	0.057
1391	M25315	Homo sapiens (clone pAT 464) potential	a \$
·		lymphokine/cytokine mRNA, complete cds.	0.057
1302	ADVIIIZI	Homo sapiens mRNA for KIAA0549 protein, partial cds	0.055
er-commander		Homo sapiens phosphatidylinositol 3-kinase, class 3	
1202	ND 4 000 647 1	(PIK3C3) mRNA > :: emb Z46973 HSPITR1 H.sapiens	
	NM_002647.1	mRNA for phosphatidylinositol 3-kinase	0.051
1384	Z18274	Homo sapiens satellite DNA	0.05
1205	7.0.50.44	Homo sapiens integral nuclear envelope inner membrane	***************************************
	L25941	protein (LBR) gene, complete cds.	0.049
	U28171	Trypanosoma cruzi kinetoplast putative maxicircle divergent	0.049
1387	X76168	R.norvegicus mRNA for connexin 30.3	0.049
	· · · · · · · · · · · · · · · · · · ·	Morganella morganii xylitol repressor, complete cds; xylitol	***************************************
		dehydrogenase, complete cds; xylulokinase, 5' end.	

1422 U32446

Table 2A Nearest Neighbor (BlastN vs. GenBank) SEO ID ACCESSION **DESCRIPTION P VALUE** Human Ig germline kappa L-chain V-region gene germline 1389 K01322 kappa L-chain V-region gene (HK189), V-kappa-1. 0.049 capsid protein {RNA 3} [tomato aspermy virus TAV-B, 1390 S72468 Blencowe, Genomic RNA, 2213 ntl 0.049 Zea mays auxin-binding protein (abp4) gene, exons 1-5 and 1391 L08426 complete cds. 0.048 1392 X96972 D.buzzatii copia element DNA 0.048 1393 X17147 Canine mRNA for thyrotropin (TSH) receptor variant 0.048 1394 L13164 Saccharomyces cerevisiae k9 killer toxin resistant 0.047 E.octocarinatus RPA2 gene for RNA polymerase I second 1395 X66451 largest subunit 0.047 1396 X75426 G.biloba (developing endosperm) ginnacin mRNA 0.047 Aedes aegypti LINE retrotransposon Juan-A including DNA binding protein and reverse transcriptase-like protein 1397 M95171 mRNA, complete coding regions. 0.047 1398 Z49076 R.prowazekii gene (unknown) 0.047 Human Ig germline kappa L-chain V-region gene germline 1399 K01323 immunoglobulin heavy chain, kappa chain, 0.047 Homo sapiens galactose-1-phosphate uridyl transferase 1400 L48713 (GALT) mutant V44L gene, exon 7 (M96246 bases 303-0.047 1401 U77310 Drosophila melanogaster porcupine mRNA, complete cds 0.047 1402 J01323 Yeast (S. cerevisiae) enolase gene (clone peno8) and flanks. 0.047 Bovine herpesvirus type 1 immedidate-early transcriptional 1403 L14321 control protein (BICP4) gene, 5' end. 0.047 Homo sapiens myotonic dystrophy-associated protein kinase 1404 L19266 and 59 genes. 0.047 1405 M58600 Human heparin cofactor II (HCF2) gene, exons 1 through 5. 0.046 1406 D86964 Human mRNA for KIAA0209 gene, partial cds 0.046 1407 Z83329 S.salar mRNA for transport associated protein Tap2B 0.046 Glyphinaphis bambusae mitochondrial cytochrome oxidase 1408 L27331 subunit I gene, 3' end, and cytochrome oxidase subunit II 0.046 Human interleukin-2 receptor alpha chain (IL2RA) gene, 1409 U57613 promoter region 0.046 Solanum tuberosum sucrose synthase gene, clone gPOSS65, 1410 U24088 complete cds. 0.046 1411 V01087 Hemagglutinin gene of influenza virus strain 0.046 1412 S76792 OX40=cell surface antigen [human, mRNA Partial, 1034 nt] 0.046 Lycopersicon esculentum class II small heat shock protein 1413 U72396 Le-HSP17.6 mRNA, complete cds 0.046 Human non-histone chromatin protein HMG1 (HMG1) gene, 1414 U51677 complete cds 0.046 1415 X98743 H.sapiens mRNA for RNA helicase (Myc-regulated dead 0.046 1416 M63868 C.hircus alpha-lactalbumin gene, exons 1-4. 0.046 Caenorhabditis elegans cosmid H06O01, complete sequence 1417 Z92970 [Caenorhabditis elegans] 0.045 Human leukocyte adhesion protein p150,95 alpha subunit 1418 M29482 gene, exons 2 - 6. 0.045 1419 Z78942 H.sapiens flow-sorted chromosome 6 HindIII fragment, 0.045 1420 L06863 Cricetulus griseus type VII collagen mRNA, 3' end. 0.045 1421 X86449 E.caballus DNA segment containing microsatellite 0.045

protein homolog (Brca1) mRNA, complete cds.

Mus musculus putative breast/ovarian cancer susceptibility

0.045

EQ II	ACCESSION	DESCRIPTION	P VALU
		Human Ig germline kappa L-chain V-region gene germline	****
142	3 K01323	immunoglobulin heavy chain, kappa chain,	0.045
		Human Ia-associated invariant gamma-chain gene, exon 1,	
142	4 M13555	clones lambda-y(1,2,3).	0.045
		Rattus norvegicus chaperonin 60 (Hsp60) and chaperonin 1	0
142	5 U68562	(CPN10) genes, nuclear genes encoding mitochondrial	0.045
		Homo sapiens DNA for choline kinase like protein and	0.073
142	6, AB003286	muscle type carnitine palmitoyltransferase I, partial and	0.045
142	7 U34372	Human tyrosine kinase TXK (txk) gene, exon 6.	0.045
	***************************************	Human non-histone chromatin protein HMG1 (HMG1) gene	0.043
142	8 U51677	complete cds	0.045
***************************************		Dictyostelium discoideum phosphatidylinositol-4,5-	10.043
1429	9 U23476	diphosphate 3-kinase (PIK1) mRNA, complete cds.	0.045
1430	Z98975	S.pombe chromosome I cosmid c19E9	0.043
********	1 X16465	Trypanosoma brucei mRNA for cysteine proteinase	0.044
Militaria in accessoration	-d:	Macaca fascicularis mitochondrial DNA for NADH	0.044
1432	D85274	dehydrogenase subunit 4, subunit 5, partial cds	0.044
	3 X16876	Soybean ENOD2B gene for Ngm-75	0.044
	U19755	Mus domesticus thyroid transcription factor 1 gene,	0.044
	L77700	Gallus gallus 18C15 mRNA, complete cds.	0.044
	AF019981	Dictyostelium discoideum HelE (helE) gene, partial cds	0.044
	7 L13469	Saccharomyces cerevisiae antiviral protein Ski2p	0.044
	3 M26238	D discoideum apora cost matein SD70	0.044
**********	U65391	D.discoideum spore coat protein SP70 gene, complete cds.	0.044
		Lycopersicon esculentum PRF (Prf) gene, complete cds Mus musculus nuclear receptor coactivator protein 2 mRNA	0.044
1440	AF000582	complete cds	3
***********	X98880	C.albicans ARG5,6 gene	0.044
	D89609		0.044
	Y13544	Oryzias latipes mRNA for choriogenin H, complete cds Homo sapiens cosmid C1	0.044
The second secon	Y14952		0.044
	114732	Mus musculus gene encoding immunoglobulin J chain	0.043
1445	Z69660	Caenorhabditis elegans cosmid F39B1, complete sequence	
	X53404	[Caenorhabditis elegans]	0.043
	U48271	Glycine max glycinin A(1a)B(1b) and A(2)B(1a) boundary	0.043
177/	U402/1	Dictyostelium discoideum UbpA deubiquitinase mRNA,	0.043
		Saccharomyces cerevisiae origin recognition complex,	a.i.iiddoondoo
		subunit 5 (ORC5) gene, complete cds > :: gb I32734 I32734	94
1//0	U24187	Sequence 9 from patent US 5589341 > :: gb I38710 I38710	5
*************	Z28177	Sequence 9 from patent US 5614618	0.043
1449		S.cerevisiae chromosome XI reading frame ORF YKL178c	0.043
1450	V05051	Drosophila melanogaster calmodulin gene exon 4 and intron	
	X05951		0.043
	L12999	Daltonia sp. mitochondrial 16S ribosomal RNA	0.043
mmmmmm	U68098	Human poly(A)-binding protein (PABP) gene, exons 6 and 7	0.043
······································	U55774	Caenorhabditis elegans cosmid F35G8.	0.043
······································		H.sapiens DNA for alu repeats	0.043
······································		Caenorhabditis elegans cosmid ZK1131.	0.043
1456	M30168	D.melanogaster nested repetitive sequences F and G,	0.043
		Human immunodeficiency virus type 2 isolate HIV2CBL21	······································
~~~~	U05350 (	gp160 envelope (env) gene, complete cds.	0.042
	U44129	Rattus norvegicus p58 mRNA, complete cds	0.042
	L13377	Staphylococcus aureus enterotoxin gene, 3' end.	0.042
1 1 / 0	X83758	D C 1 :	0.042

SEQ ID	ACCESSION	DESCRIPTION	DAZATE
		Anagrapha falcifera nuclear polyhedrosis virus protein-	PVALU
	ever ever	typrosine phosphatase, lef-2, polyhedrin, vp78 and protein	0.000
1461	U64896	kinase genes, complete cds	0.040
~~~~~	J02175	Influenza A/wsn/33 (h1n1), defective interfering 13, cdna.	0.042
		Arabidopsis thaliana DNA for inorganic phosphate	0.042
1463	D86608	transporter, complete cds	
	M93716	Pan paniscus DNA fragment.	0.042
2101	111/00/110	Astasia longa chloroplast rps7 and tufA genes for ribosomal	0.042
1465	X14385	protein S7 and elongation factor Tu respectively	4
1.00		Saccharomyces carevisiae and achiticae (CTS1.1)	0.042
1466	M74069	Saccharomyces cerevisiae endochitinase (CTS1-1) gene, complete cds.	00000000000000000000000000000000000000
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M73257		0.042
1707	141/ <i>323</i>	C.parasitica laccase (lac-1) gene, complete cds.	0.042
1468	M31126	Human pregnancy-specific beta-1-glycoprotein (SP1)	MARKET AND A STATE OF THE STATE
	U70826	mRNA, complete cds, clone hPS2.	0.042
~~~~~~~	AF000299	Fundulus heteroclitus vitellogenin II precursor mRNA,	0.041
·····	X83390	Caenorhabditis elegans cosmid E03H12	0.041
***************************************	L12582	Albinaria coerulea complete mithochondria DNA	0.041
14/2	L1438Z	Human ornithine decarboxylase (ODC) gene, 5' flanking	0.041
1472	D00655	Rattus norvegicus mRNA for scavenger receptor class B,	BOAD Military
	D89655	complete cds	0.041
······································	Z23267	P.anserina AS1 gene, complete CDS	0.041
	M63868	C.hircus alpha-lactalbumin gene, exons 1-4.	0.041
	U56440	Human His-1 gene sequence	0.04
omenion con con con con construir de la constr	L06863	Cricetulus griseus type VII collagen mRNA, 3' end.	0.04
	U02928	Dictyostelium discoideum Ax3 Rab7 mRNA, complete cds.	0.04
	M19828	Human apolipoprotein B-100 (apoB) gene, exons 22 through	0.04
1480 1	M34434	Human angiotensin-converting enzyme (ACE) gene, 5' end.	0.04
		Human DNA sequence from cosmid L21F12, Huntington's	<u> </u>
1481 2	Z68884	Disease Region, chromosome 4p16.3	0.04
Willia novice		Pig alveolar macrophage-derived chemotactic factor-I	<u> </u>
1482 N	M99367	(AMCF-I) mRNA, complete cds.	0.04
OAKSTANIAN A		Human hindlimb expressed homeobox protein backfoot (Bft)	
1483 L	J70370	mRNA, complete cds	0.04
		Rattus norvegicus liver microsomal carboxylesterase mRNA,	
	J <b>10698</b>	complete cds.	0.04
1485 L	J <b>28171</b>	Trypanosoma cruzi kinetoplast putative maxicircle divergent	0.04
	J <b>79297</b>	Human clone 23589 mRNA sequence	0.04
1487 X	K13764	Yeast centromere CEN1 repetitive DNA PSS113, part of	0.04
1488 Z	24568	TT · / ~ ~ ~ ~ ~ ~	0.04
1489 N	<b>1</b> 63868		0.04
1490 X	X64143		0.039
1491 N			0.039
		Mus musculus breast and ovarian cancer susceptibility	V.UJJ 
1492 U	J36475	manufacture (D. 1) David	0.020
1493 X	~ ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		0.039
	The Control of the Co	Rattus norvegicus carcinoembryonic antigen-related protein	0.039
1494 M	132475	(00) (1)	0.020
1495 D	·····		0.039
1990 Martin Carris State Contract Contr	Marian Commission of the Commi	Homo saniens zinc finger protein (7-E20) DALA	0.039
1497 X	······································		0.039
1498 Z		D 1: :: -	0.039
		TT	0.038
Janone and Salara	······································		0.038
1500 X	······································	the state of the s	0.038 0.038

1521 U09865

1522 U67815

1523 AE000466

1524 M80785

1525 L37035

1526 M15009

1527 U67500

1529 X52956

1530 U80581

1531 Z69918

1532 Z98031

1533 D45858

1534 L24549

1535 U66291

1536 X63436

1537 L23498

1538 AF003086

1539 U17377

1540 L23498

1541 X85117

1528 AB000044

		BlastN vs. GenBank)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUE
1501	D89609	Oryzias latipes mRNA for choriogenin H, complete cds	0.038
1502	Z49535	S.cerevisiae chromosome X reading frame ORF YJR035w	0.038
1503	L13377	Staphylococcus aureus enterotoxin gene, 3' end.	0.035
1504	L12582	Human ornithine decarboxylase (ODC) gene, 5' flanking	0.035
	and to the transfer the transfer and the transfer transfer and the transfer and the transfer and the transfer a	Caenorhabditis elegans cosmid ZK1321, complete sequence	
1505	Z48584	[Caenorhabditis elegans]	0.023
1506	Y00204	Xenopus laevis mRNA fragment for nucleoplasmin	0.021
		light-chain fibroin [Galleria mellonella=waxmoths, larvae,	
1507	S77817	cocoons, posterior silk glands, mRNA, 1191 nt	0.021
		Trypanosoma cruzi (strain La Cruz, Jalisco) ribosomal RNA	
1508	L13926	transcribed spacer and 18S ribosomal RNA gene, 5' end.	0.02
1509	Z78910	H.sapiens flow-sorted chromosome 6 HindIII fragment,	0.02
1510	AB007976	Homo sapiens mRNA, chromosome 1 specific transcript	0.02
1511	U01307	Human scRNA (BC200 beta) pseudogene.	0.017
1512	J03998	P.falciparum glutamic acid-rich protein gnen, complete cds.	0.017
1513	X93497	H.sapiens TRAP gene, intron 4 (partial)	0.017
1514	M15656	Human aldolase B (ALDOB) gene, exons 7 through 9.	0.016
		H.sapiens ACPP gene for prostatic acid phosphatase (non-	
1515	X74965	coding region)	0.016
1516	X75653	A.longa plastid genes for ribosomal proteins and tRNAs	0.016
1517	X75653	A.longa plastid genes for ribosomal proteins and tRNAs	0.016
2000		Human MLC1emb gene for embryonic myosin alkaline light	
1518	X58852	chain, exon 2	0.016
1519	U69695	Mus musculus hyaluronan synthase homolog mRNA,	0.016
1520	S45332	erythropoietin receptor [human, placental, Genomic, 8647	0.016

Alcaligenes eutrophus pyruvate dehydrogenase

Human primary Alu transcript

complete genome

dihydrolipoamide dehydrogenase (pdhL), and ORF3 genes,

Escherichia coli K-12 MG1655 section 356 of 400 of the

M.musculus tissue factor promoter (Cf-3) gene, exon 1.

Mouse steroid 21-hydroxylase A (21-OHase A) gene,

Rhizoctonia solani 5.8S rRNA gene, complete sequence

Drosophila virilis brown protein (bw) gene, complete cds.

Methanococcus jannaschii section 42 of 150 of the complete 0.016

0.016

0.016

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0.016

	Human CAMII-psi3 calmodulin retropseudogene	0.016
	Pleurodeles waltl Wnt-7a mRNA, complete cds	0.016
	Human DNA sequence from cosmid 91K3, Huntington's	
	Disease Region, chromosome 4p16.3 contains CpG island	0.016
	Human immunodeficiency virus type 1 nef gene	
	immunodeficiency virus type 1 nef gene (strain KU15-1)	0.016
	Mus musculus mRNA for synaptotagumin III, complete cds	0.016
	Gallus gallus Gi2 protein alpha-subunit mRNA, complete	0.016
	Catagonus wagneri cytochrome b gene, mitochondrial gene	
	encoding mitochondrial protein, partial cds	0.016
	B.taurus mRNA for poly(A) polymerase	0.016
	Bovine microsatellite repeats	0.015
	Plasmodium falciparum transcription factor homolog	
_	PfSNF2L mRNA, complete cds	0.015
	Strongylocentrotus purpuratus cortical granule protein with	
	LDL-receptor-like repeats mRNA, partial cds.	0.015
_	Bovine microsatellite repeats	0.015
	H.sapiens epb72 gene exons 2,3,4,5,6,7	0.015
	T 11 04	
	Table 2A	
	Page 38 of 61	

SEQ ID	ACCESSION	DESCRIPTION	P VALU
	Z16906	H. sapiens (D14S73) DNA segment containing	0.015
CONTRACTOR	X99400	S.pneumoniae dacA gene and ORF	0.015
	M32061	Rat alpha-2B-adrenergic receptor (RNG-alpha-2) mRNA,	0.015
	X80930	S.cerevisiae RHC18 genes	0.015
1343	A00930	Alcaligenes eutrophus pyruvate dehydrogenase	10.015
1546	U09865	dihydrolipoamide dehydrogenase (pdhL), and ORF3 genes,	0.015
1340	007803	Mus musculus BALB/c of p65 gene encoding p65 subunit of	
1547	Z22952	transcription factor NF-kappaB	0.015
1 347	LLL93L	Mus musculus parathyroid hormone/parathyroid hormone	0.013
1548	L34610	related-peptide receptor (PTHR) gene, exons 5 - 9.	0.015
1340	LJ4010	H.sapiens CpG island DNA genomic Mse1 fragment, clone	0.013
1540	Z54850	169b5, reverse read cpg169b5.rt1a	0.015
0x7494040404047434040447 <b>4</b> 470	U01307	Human scRNA (BC200 beta) pseudogene.	0.015
	M54994	Human bile salt-activated lipase (BAL) mRNA, complete	0.015
1551	WI34994	Human immunodeficiency virus type 1 nef gene	0.013
1552	709021		0.015
January Strategy and Company of the	Z98031 Z21858	immunodeficiency virus type 1 nef gene (strain KU15-1)  M.Musculus mRNA for P domain protein	0.015
		Human HLA-B-associated transcript 2 (BAT2) gene, 3' end.	kan marka kan marka kan kan kan kan kan kan kan kan kan k
	M33518 AB001383	Rattus norvegicus mRNA for sialoprotein, complete cds	0.015
1333	ABUU1383	Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	0.015
1556	U72787		0.014
1330	U12181	complete sequence [Homo sapiens] Sambucus nigra ribosome inactivating protein precursor	(U.U14
1557	AE012900	mRNA, complete cds	0.014
	AF012899 J03764		0.014
1000	JUS / 04	Sambucus nigra ribosome inactivating protein precursor	10.014
1550	AF012899	mRNA, complete cds	0.014
1339	AFU12099	Fruitfly strain g20 mitochondrial DNA, A+T-rich region,	0.014
1560	AB003097	partial sequence	0.014
	AC001017	Homo sapiens (subclone 2 g8 from P1 H43) DNA sequence	
	Z48484	H.sapiens gene for tissue-type plasminogen activator	0.014
	AC001460	Homo sapiens (subclone 2 f4 from BAC H107) DNA	0.014
1303	AC001400	Alcaligenes eutrophus pyruvate dehydrogenase	}
1564	U09865	dihydrolipoamide dehydrogenase (pdhL), and ORF3 genes,	0.014
1304	009003	Dictyostelium discoideum plasmid Ddp1 D2 orf, D1/D3 orf,	0.014
1565	U00691	G4/D5 orf, G5/D6 orf, G1 orf, G2/G3/D4 orf, complete cds	0.014
	D16482	Sarcophaga peregrina mRNA for poly(ADP-ribose)	0.014
1300	D10402	Sambucus nigra ribosome inactivating protein precursor	0.014
1567	AF012899	mRNA, complete cds	0.014
upunununun perununun pununun k	M63599	Human myelin basic protein (MBP) gene, exon 1.	0.014
	U03891	Human phorbolin I mRNA, partial cds.	0.014
a, contra de la contractica del la contractica del la contractica de la contractica	X53422	D. grimshawi s18, s15, s19 and s16 chorion protein genes	0.014
13/0	A33444 	Yeast mitochondrial ori(o) repeat unit of petite mutant 4	0.013
1571	X01870		0.013
13/1	AU10/U	Caenorhabditis elegans cosmid K05D4, complete sequence	0.013
1572	702004		0.012
market construction and the second contract of the second contract o	Z92804 M76377	[[Caenorhabditis elegans]	0.013
	M76377 D29801	Human cysteine-rich protein (CRP) gene, exons 3 and 4.	0.013
marrow and a second		Mouse mRNA for unknown product, complete cds	0.013
13/3	U77984	Sus scrofa microsatellite S0058 sequence	0.013
1576	755000	H.sapiens CpG island DNA genomic Mse1 fragment, clone	0.012
13/6	Z55882	70g8, reverse read cpg70g8.rt1a	0.013
1 - 77	3700521	Human interferon genes LeIF-L and LeIF-J and pseudogene	0.013
13//	V00531	LeIF-M with intergenic regions. These genes are located on S.cerevisiae chromosome IV reading frame ORF YDR065w	0.013

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
ord m	ACCESSION	Herpes simplex virus type 1 (HSV-1) genome, rightmost part	
1570	D00374	of the long unique region (UL) and all of the internal long	0.013
			~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
	U53502	Arabidopsis thaliana chromosome I cosmid g17311 DNA.	0.013
1381	K03196	Human interferon-beta-3 gene.	0.013
1500	) (02.45C	Zea mays (clone pCIB808) encoding maize nitrite reductase	0.013
1382	M23456	(NiR) cDNA to mRNA, partial cds. H.sapiens F8VWFL DNA of the von Willebrand factor	0.013
1500	V((120		0.012
***************************************	X66120	pseudogene (5'portion)	0.013
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M63962	Human gastric H,K-ATPase catalytic subunit gene, complete	elynoodikkiiliitoon oneliisiiristaan eesistai kannee
emenous extremos conservations of	AF039592	Homo sapiens succinate dehydrogenase subunit C	0.007
····	Y10908	B.cereus cysA and wapA genes	0.007
	AF048988	Homo sapiens MutS homolog 5 (MSH5) gene, exons 1	0.007
~~~~~~~~~~	AF048988	Homo sapiens MutS homolog 5 (MSH5) gene, exons 1	0.007
	D90822	E.coli genomic DNA, Kohara clone #331(40.1-40.4 min.)	0.007
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	D87903	Mouse mRNA for ARF6, complete cds	0.006
1591	L38851	Mycobacterium tuberculosis cell surface protein	0.006
		Human neuronal nitric oxide synthase (nNOS) gene,	
	U66362	alternatively spliced 5' exon Tex 2 and flanking sequences	0.005
	L11670	Human transmembrane glycoprotein (CD53) gene, exons 2	0.005
	U20587	Sus scrofa domestica sister of P-glycoprotein	0.005
	U53016	Human DXYS154 microsatellite sequence	0.005
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z29641	Zea mays of USE gene encoding U3snRNA	0.005
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	D16473	Human mRNA, Xq terminal portion	0.005
manuscus various menter mili	X02175	Schizosaccharomyces pombe cdc10 start gene	0.005
1599	U25029	Human glucocorticoid receptor alpha mRNA, variant 3'	0.005
		Human DNA sequence from cosmid N2E9 on chromosome	
···	Z68685	22, complete sequence [Homo sapiens]	0.005
and the second s	U40369	Human spermidine/spermine N1-acetyltransferase	0.005
1602	U06965	Aphytis melinus mitochondrion 16S rRNA gene, partial	0.005
1.00	704044	Human DNA sequence from cosmid U65A4, between	
·	Z81014	markers DXS366 and DXS87 on chromosome X *	0.005
in a consequence of	X91923	<u> </u>	0.005
	Z29641	Zea mays of USE gene encoding U3snRNA	0.005
	L11670	Human transmembrane glycoprotein (CD53) gene, exons 2	0.005
~~~~~	U15605	Nicotiana glutinosa virus resistance (N) gene, complete cds.	0.005
	X57698	ทั้งเสียนาากราบน เพรียนสมาร์ที่สมาร์ที่สมาร์ที่สมาร์ที่ พระพระบบเมื่อการ์ที่สมาร์ที่มากระบบเมื่อการ์ที่สมาร์ที	0.005
	L81391	· · · · · · · · · · · · · · · · · · ·	0.005
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	X81789	H.sapiens mRNA for splicing factor SF3a60	0.005
	X82818	H.sapiens PTP1C/HCP gene	0.005
errouewithtusseuwww.ve.noneouily	U33465	Drosophila melanogaster S element and flanking sequence,	0.005
1613	Z75042	S.cerevisiae chromosome XV reading frame ORF YOR134w	0.005
		Mouse T cell receptor rearranged alphachain variable	
·····	M98574	region, N_region, joining region, and constant region	0.005
1615	X51907		0.005
userendan		Human retinoic acid-responsive protein (NN8-4AG) mRNA,	
1616	U50383	complete cds	0.005
tenne, warr		Human cardiac alpha-myosin heavy chain (MYH6) gene,	
Nikanoonii (Nikanoo ooni) Nikanoolija	M25140	£	0.005
~~~~~~	X66062	G.max tufA gene for chloroplast translation elongation	0.005
1619	U16345		0.005
Webserman		Sambucus nigra ribosome inactivating protein precursor	
1 (20)	AF012899	mRNA, complete cds	0.005

1648 U08421

1649 Z17089

1650 L35531

1651 M55120

1652 J00922

1653 U69695

1654 L15248

1655 L35676

1656 L48612

1657 Y08925

1658 M60441

1659 X91857

Table 2A Nearest Neighbor (BlastN vs. GenBank) SEQ ID ACCESSION DESCRIPTION P VALUE Saccharomyces cerevisiae chromosome IV lambda 3073 and 1621 U43834 flanking region extending into right telomere 0.005 Caldocellum saccharolyticum celB gene for cellobiohydrolase/endocellulase 1622 X13602 0.005 1623 J04809 Human cytosolic adenylate kinase (AK1) gene, complete 0.005 Homo sapiens neuroendocrine-specific protein (NSP) gene, 1624 L49144 exons 1B and 3 0.005 Plasmodium falciparum (clone HB3) heat shock protein 86 1625 L34028 gene, complete cds. 0.005 Human retinoic acid-responsive protein (NN8-4AG) mRNA, 1626 U50383 0.005 complete cds Lucilia cuprina peritrophin-95 precursor gene, partial cds. 1627 U23829 0.005 Helicobacter pylori 26695 section 107 of 134 of the 1628 AE000629.1 0.005 Human adenine phosphoribosyltransferase (APRT) gene, 1629 M16446 complete cds. 0.005 Human retinoic acid- and interferon-inducible 58K protein 1630 U34605 RI58 mRNA, complete cds. 0.005 Human cardiac alpha-myosin heavy chain (MYH6) gene, 1631 M25140 exons 2, 3 and 4. 0.005 Mouse MHC (Qa) Q10-k gene for class I antigen 1632 X16426 0.005 1633 M83985 Mouse phosphoprotein (F1-20) mRNA, complete cds. 0.005 1634 U31850 Human dystonin isoform 1 mRNA, partial cds 0.005 1635 U31850 Human dystonin isoform 1 mRNA, partial cds 0.005 O.berteriana chloroplast ORF80 (exon 1 and 2) and 1636 X64615 ORF2280 sequences 0.005 1637 X52647 E.coli dbpA gene for DEAD box protein A 0.005 1638 D90773 E.coli genomic DNA, Kohara clone #262(30.3-30.5 min.) 0.005 S.glaucescens novel deletion/rearrangement sequence, 1639 M62946 partial sequence. 0.005 1640 M88597 Saccharomyces cerevisiae STP1 gene, complete cds. 0.005 Homo sapiens (clone HG52) Z-crystallin/quinone reductase (CRYZ) gene sequence. 1641 L31521 0.005 1642 D79986 Human mRNA for KIAA0164 gene, complete cds 0.004 1643 AC002183 Homo sapiens (subclone 2 h8 from BAC H111) DNA 0.004 1644 Z29641 Zea mays of USE gene encoding U3snRNA 0.004 1645 L22415 Homo sapiens DNA sequence, repeat region. 0.004 Chlamydomonas reinhardtii chloroplast 30S ribosomal 1646 U17357 protein S4 (rps4) gene, complete cds. 0.004 Rat 3-beta-hydroxysteroid dehydrogenase/delta-5-delta-4-1647 M67465 ene-isomerase mRNA, complete cds. 0.004

P.falciparum aarp3 gene, exon	
Sheep lambda immunoglobulin V gene	·.
Piromyces sp. mRNA for mannanase A	
Table 2A	
Page 41 of 61	
<u> </u>	

Murine rotavirus EC outer capsid protein VP4 gene,

Human cystic fibrosis transmembrane conductance regulator

Homo sapiens (subclone H8 2 e7 from P1 35 H5 C8) DNA

Theileria parva (clone pTprUgB) ORF genes, partial cds.

H. sapiens (D3S1309) DNA segment containing

Gallus gallus ovalbumin (Y) gene, complete cds.

Human (clone: pHyTM1/60(R)) DNA sequence.

Mus musculus hyaluronan synthase homolog mRNA,

Human Alu repeat region.

(CFTR) gene, exon 14b

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SEQ ID	ACCESSION	DESCRIPTION	P VALU
	Z35948	S.cerevisiae chromosome II reading frame ORF YBR079c	0.002
*****	X16277	Human gene for ornithine decarboxylase ODC (EC 4.1.1.17)	
	X78608	G.gallus genomic DNA repeat region, clone 9C2	0.002
	U48449	Human skeletal muscle ryanodine receptor gene	0.002
1003	V40447	Human breakpoint in translocation V-kappa gene region	(U.UUZ
1661	X51875	(WB) (partial) (537 bp)	0.002
management and	Z24205	H. sapiens (D12S348) DNA segment containing	ágarana a sa
1003	2,24203	Mus musculus cartilage-derived retinoic acid-sensitive	0.002
1666	1106613		0.002
	U03012	31 3 7 1	0.002
1667	N/C1020	Human alpha-1,3-mannosyl-glycoprotein beta-1, 2-N-	0.000
1007	M61829	acetylglucosaminyltransferase (MGAT) gene, complete cds.	0.002
1.660	14/1000	Human alpha-1,3-mannosyl-glycoprotein beta-1, 2-N-	0.000
and the second second second second second	M61829	akananan kan Kamanan manan kan mengan mengan mengendaran kan menan bebaak mengan melang mengentian mengan meng	0.002
~~~	V00571	Human gene encoding prepro form of corticotropin releasing	
and the second second second	D29760	Candida maltosa gene for chitin synthase 1, partial sequence	
	AJ001817	Dama sp. mRNA for bone morphogenetic protein 2	0.002
	Z23575	H. sapiens (D17S926) DNA segment containing	0.002
	X59359	T.marmorata mRNA for acetylcholinesterase	0.002
1674	D12519	Rat SAP gene for synaptotagmin associated 35kDa protein	0.002
		Mus musculus glucose-6-phosphate dehydrogenase protein,	C. Constitution of the Con
	U88534	exons 10, 11 and partial cds	0.002
	Z24391	H. sapiens (D11S1350) DNA segment containing	0.002
	M31773		0.002
1678	U28014	Human cysteine protease (ICErel-II) mRNA, complete cds.	0.002
1679	M33518	Human HLA-B-associated transcript 2 (BAT2) gene, 3' end.	0.002
		Pseudomonas aeruginosa heptosyl transferase II	
***************************************	U70983	······································	0.002
	Z74854	S.cerevisiae chromosome XV reading frame ORF YOL112w	
1682	M13498		0.002
-		Streptococcus pyogenes putative multiple membrane domain	
aan ee aan aan aan aan aan ah	U17382	protein gene, complete cds.	0.002
1684	U14189	Plasmodium falciparum MCP1 mRNA, complete cds	0.001
-		Saccharomyces cerevisiae mitochondrial petite mutant BB5	
1685	M27314	origin of replication DNA.	0.001
	M64089	Dictyostelium discoideum calmodulin (calA) gene, complete	0.001
1687	J05138	Rabbit calcium binding protein (calreticulin) mRNA,	0.0009
***************************************		Human clone 199288 defective mariner transposon Hsmar2	
1688	U92017	mRNA sequence	0.0008
		Homo sapiens mRNA for synaptosome associated protein of	***************************************
1689	AJ011915	23 kilodaltons, isoform A	0.0008
1690	AJ223734	Sus scrofa SCAMP1 gene, exon 1 and joined CDS	0.0008
1691	Y08924	P.falciparum mRNA for AARP2 protein	0.0006
	X58139	Human coxVIb gene, last exon and flanking sequence	0.0006
and the second second	U47853	Araneus diadematus fibroin-1 (ADF-1) mRNA, partial cds	0.0006
naa, ta gadhaagan iya iy gadhaa na iya iya gadhaa ba	i engonicario den contintia e contrata entrata describer en el consociedad de consociedad de consociedad de co	Plasmodium falciparum (clone Dd2) heat shock protein 86	ga tirraring kanamang panah-hang pilinama,
1694	L34027	1	0.0006
anna ann ann ann ann ann ann ann ann an	D88271	Contraction of the contraction o	0.0006
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	AC001546		0.0006
		H.sapiens telomeric DNA sequence, clone 16QTEL024,	***************************************
1697	Z96325	,	0.0006
		Saccharomyces cerevisiae Nmd2p (NMD2) gene, complete	
1698	U14974		0.0006
· marante marante marante marante marante	AB002331	бростический министический испаньит чести и при при при при при при при при при	0.0006

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
ing a state of the same of the		Escherichia coli K-12 MG1655 section 354 of 400 of the	Y VILLO
1700	AE000464	complete genome	0.0006
	Z79060	H.sapiens flow-sorted chromosome 6 HindIII fragment,	0.0005
1701	<i>L</i> 19000	H.sapiens CpG island DNA genomic Mse1 fragment, clone	0.0003
1702	Z60233	197c9, reverse read cpg197c9.rt1a	0.0005
	U15018	Dugbe virus L protein gene, complete cds	a grant and the second
1703	U13010	H. sapiens genomic DNA (leukocyte), corresponding to the	0.0005
1704	V77607	,	0.0005
unicerretiscorpetiscorpu petitica, ult	X77607	integration site of HPV 6a DNA in a tonsillar carcinoma	0.0005
	M59428	T.thermophila ribosomal protein L37 (L37) mRNA,	0.0005
	M59428	T.thermophila ribosomal protein L37 (L37) mRNA,	0.0005
	AC002219	Homo sapiens (subclone 2_d11 from P1 H43) DNA	0.0005
and the second second second second	X95276	P. falciparum complete gene map of plastid-like DNA (IR-B)	
AND THE RESIDENCE AND THE PARTY OF THE PARTY	L18972	Homo sapiens anonymous gene, complete cds	0.0005
1/10	U96974	Homo sapiens MET proto-oncogene, intron 5, 3' end	0.0005
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	and the second s
	Z60916	39a5, forward read cpg39a5.ft1c	0.0005
	X99587	A.brasilense ipdC, gltX & cysS genes	0.0005
1713	J03998	P.falciparum glutamic acid-rich protein gnen, complete cds.	0.0005
		Escherichia coli K-12 MG1655 section 354 of 400 of the	300 American
	AE000464	complete genome	0.0005
1715	U21730	Human 5'-nucleotidase (CD73) gene, partial cds.	0.0004
and the second second second second	M22970	Human pancreatic phospholipase A-2 (PLA-2) gene, exons 1	0.0003
1717	U37022	Human cyclin-dependent kinase 4 (CDK4) gene, complete	0.0002
1718	AC001517	Homo sapiens (subclone 1_g5 from P1 H49) DNA sequence	0.0002
1719	X90383	A.thaliana DNA for Y13 gene	0.0002
		Homo sapiens zinc finger protein homologous to Zfp-36 in	
Í		mouse (ZFP36) mRNA > :: gb M92843 HUMG0S24A	COMPANY
1720	NM_003407.1	H.sapiens zinc finger transcriptional regulator mRNA,	0.0002
1721	M86528	Human neurotrophin-4 (NT-4) gene, complete cds.	0.0002
1722	M86528	Human neurotrophin-4 (NT-4) gene, complete cds.	0.0002
1723	U19241	Homo sapiens interferon-gamma receptor alpha chain gene,	0.0002
		Human thymidine kinase gene, complete cds, with clustered	
1724	M15205	Alu repeats in the introns.	0.0002
		Mouse Y specific region of AC11 DNA sequence, LINE	
1725	M87510	repeat and Bkm satellite.	0.0002
		Human DNA sequence from phage LAW2 from a contig	
		from the tip of the short arm of chromosome 16, spanning	
1726	Z84723	2Mb of 16p13.3 Contains Interleukin 9 receptor pseudogene	0.0002
on at the common personness and process and p	X01392	Human apolipoprotein CIII gene and apo AI-apo CIII	0.0002
and and an arrange and a second of the secon	Z92910	H.sapiens HFE (HLA-H) gene	0.0002
AND A CONTRACTOR CONTRACTOR AND PROCESSION OF	D87001	Human (lambda) DNA for immunoglobulin light chain	0.0002
		Yeast (S.cerevisiae) mitochondrial autonomously replicating	
1730	M35612	sequence DNA.	0.0002
reserve en	Z16956	H. sapiens (D2S154) DNA segment containing	0.0002
	L42456	Mus musculus TGF-1 gene, promoter region and exon 1.	0.0002
	X90383	A.thaliana DNA for Y13 gene	0.0002
	L35657	Homo sapiens (subclone H8 5 a10 from P1 35 H5 C8) DNA	
~~~	M57902		
1/33	1 <b>41</b> 3 / 3U4 	Mouse transforming growth factor-beta-1	0.0002
1726	1167167	Homo sapiens intestinal mucin (MUC2) gene, promoter	0.0000
1/30	U67167	region and partial cds	0.0002
		Homo sapiens parathymosin (PTMS) mRNA > ::	
1777	NM_002824.1	gb M24398 HUMTHYP Human parathymosin mRNA,	0.0002

SEQ ID	ACCESSION	DESCRIPTION	P VALU
SEQ ID	ACCESSION	HIV-1 patient JO population variant JOS17 from USA,	II VALO
1720	U96566	envelope glycoprotein, C2-V5 region (env) gene, partial cds	0.0003
	AF043461		9e-005
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Homo sapiens I-REL gene, exon 9	\$
	M96943	Human profilaggrin gene exons 1-3, 5' end.	8e-005
	S45332	erythropoietin receptor [human, placental, Genomic, 8647	7e-005
·	S45332	erythropoietin receptor [human, placental, Genomic, 8647	7e-005
····	L28125	Podospora anserina beta transducin-like protein	7e-005
	U53017	Human microsatellite marker sJCW13	6e-005
erromania in a company and a c	M86528	Human neurotrophin-4 (NT-4) gene, complete cds.	6e-005
announce and the second	D17554	Human mRNA for DNA-binding protein, TAXREB107,	6e-005
1748	U26556	Human ferritin H (FTHL13) pseudogene.	6e-005
1749	Z79060	H.sapiens flow-sorted chromosome 6 HindIII fragment,	6e-005
1750	AC001460	Homo sapiens (subclone 2_f4 from BAC H107) DNA	6e-005
1751	AC001033	Homo sapiens (subclone 2_c2 from P1 H48) DNA sequence	6e-005
······································	<b>34-18-18-18-18-18-18-18-18-18-18-18-18-18-</b>	Homo sapiens MHC class II HLA-DRB1 (HLA-DRB1*10)	·
1752	AF007883	intron 1 sequence	6e-005
umanamanan manan mumanan fi	U12971	Tetrahymena thermophila CU428.1VII micronuclear M	6e-005
·	X83872	H.vulgaris mRNA for cAMP response element binding	6e-005
		Human DNA sequence from cosmid 92M18, BRCA2 gene	
1755	Z73360	region chromosome 13q12-13	5e-005
	X85116	H.sapiens epb72 gene exon 1	5e-005
<del>~~~</del> ~~~~~	L81639	Homo sapiens (subclone 2_b2 from P1 H39) DNA sequence	3e-005
nichteren er eine er e	D16184		3e-005
and the second second		Chicken mRNA for nuclear b-Zip protein MafF, complete	
<i>เพรา</i> ยความสมาชากสาขากสมาชาก ค.ศ. พ.ศ. พ.ศ. พ.ศ. พ.ศ. พ.ศ. พ.ศ. พ.ศ.	AF100694	Mus musculus Pontin52 mRNA, complete cds	3e-005
	Z30978	G.gorilla DNA for Mhc Alu elements	3e-005
1761	L23429	Canis beta-galactosides-binding lectin (LGALS3) mRNA,	2e-005
4 = 60	~~~	Human interferon-inducible gene IFI-56K 5' region (56 kDa	
umanocome en mangapananció	X06559	coding capacity of unknown function)	2e-005
	AC001546		2e-005
	L35658	Homo sapiens (subclone H8 9_d12 from P1 35 H5 C8)	2e-005
å	D16472	Human mRNA, Xq terminal portion	2e-005
1766	M83665	Human high mobility group 2 protein (HMG-2) gene,	2e-005
		Human T cell receptor beta (TCRBV9S1, TCRBV7S1)	STATE OF THE PARTY
	U07977	genes, TCRBV inserted and TCRBV deleted haplotype,	2e-005
1768	D83227	Populus nigra gene for extensin like protein, complete cds	2e-005
1769	U42053	Mustela vison microsatellite repeat (Mvi 24).	2e-005
	CONTRACTOR AND AND CONTRACTOR CON	Homo sapiens Na+/glucose cotransporter (SGLT1) gene,	
1770	L29339	exon 15 and complete cds.	2e-005
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z33620	M.musculus (Balb/c) GATA-3 gene (partial)	2e-005
	D88271	Human (lambda) DNA for immunogloblin light chain	2e-005
·····		Homo sapiens DNA from chromosome 19-cosmid f19399	e acresina arrivana arrivana e a a a a a a a a a a a a a a a a a
1773	AD000833	-	2e-005
	U36755	Human thrombin receptor (F2R) gene, 5' region and partial	2e-005
and the second s	AC002252	Homo sapiens (subclone 1 g7 from BAC H76) DNA	2e-005
		Escherichia coli K-12 MG1655 section 354 of 400 of the	26-003
1776	AE000464	complete genome	20.005
ويأسيس فالمراس ومواده والموجود وموادي المراس			2e-005
omenne and a second and a second dis-	X94768	A	1e-005
and the second s	U57058	Human WD protein IR10 pre-mRNA, partial cds	9e-006
nishistraman na indinana na ana indinana na ana indinana na ana indinana indinana indinana indinana indinana i	AC001603	Homo sapiens (subclone 2_a9 from PAC H92) DNA	8e-006
	747046	Human cosmid QLL2C9 from Xq28	7e-006
1780	nage public that is a part of the design of		,,
1781	U93275 X60653	Mus musculus glucokinase gene, 5' flanking region	7e-006 7e-006

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
CONTRACTOR OF THE PROPERTY OF	L13381	Plasmodium falciparum HB3\W2 transport protein	6e-006
		Homo sapiens TRE17 oncogene-associated G0S19-	
1785	U97576	2/MIP1alpha gene, downstream sequence	6e-006
economic de la company de la c	Y11204	V.carteri gene encoding volvoxopsin	6e-006
		H. sapiens (D4S409) DNA segment containing (CA) repeat;	(
1787	Z16794	clone AFM183xd6; single read	6e-006
Water and the constitution of the constitution	D83737	Human coagulation factor XII gene, intron 2	6e-006
*******	X04871	Paramecium primaurelia macronuclear DNA telomeric	6e-006
1102		Human L1Heg repetitive element from the intergenic region	00-000
1790	M14292	of the epsilon and G-gamma globin genes.	6e-006
1770	1411-72/2	Homo sapiens amine oxidase, copper containing 3	00-000
		gb U39447 HSU39447 Human placenta copper monamine	
1701	NM 003734.1	oxidase mRNA, complete cds	6e-006
Water and the second of the se	M27147	Human alpha-2-plasmin inhibitor, allele A, 5' end.	6e-006
11/2	1712 / 1 7 /	Human DNA sequence from intron 22 of the factor VIII	00-000
		gene, Xq28. Contains the end of a 9.5kb repeated region,	
1703	X86012	int22h-1, involved in many cases of haemophilia	6e-006
*******	M33216	Human aortic-type smooth muscle alpha-actin	3e-006
·····	Z83334	H.sapiens RPS3a gene	2e-006
***************************************	M57682		2e-006
Transcription of the second	M19817	Human apolipoprotein B-100 (apoB) gene, intron J.	2e-006
	Z24068	H. sapiens (D22S427) DNA segment containing	2e-006
	Z50155	X.laevis mRNA for insulin-like growth factor I receptor	2e-006
********	Y12839	H.sapiens BH30 mRNA	2e-006
1000	112039	Human DNA sequence from cosmid U39H5, between	126-000
1801	Z70041	markers DXS6791 and DXS8038 on chromosome X	2e-006
	Z80128	H.sapiens CACNL1A4 gene, exons 16 and 17	2e-006
***************************************	U80893	Mus musculus CAG trinucleotide repeat mRNA, partial	2e-006
1005	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	H.sapiens CpG island DNA genomic Mse1 fragment, clone	ZC-000
1804	Z63192	<sup>1</sup> 7a7, forward read cpg7a7.ft1d	2e-006
Memory March	U72964	Human hepatocyte nuclear factor 4-alpha gene, exon 5	2e-006
***************************************	AC002183	Homo sapiens (subclone 2 h8 from BAC H111) DNA	2e-006
		annexin II=36 kDa calcium-dependent phospholipid-binding	20-000
1807	S73557	protein [rats, RBL-2H3 basophilic leukemia cells, mRNA,	2e-006
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U79258	Human clone 23732 mRNA, partial cds	8e-007
1000		H.sapiens CpG island DNA genomic Mse1 fragment, clone	100-007
1809	Z62146	64b2, forward read cpg64b2.ft1a	8e-007
	U44381	Human tissue inhibitor of metalloproteinases-2	8e-007
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	
1811	Z65575	47c5, reverse read cpg47c5.rt1a	7e-007
rearrangement and the contraction of the contractio	J03764	Human, plasminogen activator inhibitor-1 gene, exons 2 to	7e-007
	X75349	H.sapiens 5'flanking DNA for clotting factor IX	7e-007
		Human DNA sequence from cosmid 92M18, BRCA2 gene	/C-00/
1814	Z73360	region chromosome 13q12-13	7e-007
	27, 5 5 <b>6 6</b>	Human DNA sequence from cosmid 92M18, BRCA2 gene	70-007
1815	Z73360	region chromosome 13q12-13	7e-007
······································	X77624	H.sapiens simple sequence clone pg2m3, 5' flank and repeats	
1010	AET TOLT	SCL/TCL5/tal-1=stem-cell leukemia {germline chromosome	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
		3 translocation/deletion breakpoint} [human, bone marrow	
	046055	cells, Genomic Mutant, 239 nt]	7e-007
1217	N/46X57		
1817 1818	······································		
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	J03998	P. falciparum glutamic acid-rich protein gnen, complete cds. Sambucus nigra ribosome inactivating protein precursor	7e-007

SEQ ID	ACCESSION	DESCRIPTION	DAZATII
			P VALUI
1820	U47654	Human pyruvate kinase PK-R gene, partial cds, and pyruvat	5
1021	047004	kinase PK-L gene, complete cds.	3e-007
1821	U78096	Human macrophage colony stimulating factor receptor (c-	***************************************
Mark the Control of t	2 L76927	fms) gene, exon 1A, 2 and partial cds	3e-007
	U22086	Human galactokinase (GALK1) gene, complete cds	3e-007
	1 J03069	Ursus americanus clone G10H GT and ATTT microsatellite	3e-007
	X82640	Human MYCL2 gene, complete cds.	3e-007
CONSTRUCTION OF CONTRACTOR OF	endonesses protesses a se se consesses que proprie se consesses que partir de la consesse de la consesse de la	D.melanogaster mRNA for alpha 1,2 mannosidase	3e-007
1820	U18671	Human Stat2 gene, complete cds.	2e-007
1000	* T 0000 C	Human major breakpoint cluster region (BCR) gene, exons	1 🛊
~~~~~	L02935	3 and repeat regions.	2e-007
*************	L04193	Human lens membrane protein (mp19) gene, exon 11.	2e-007
1829	AC001050	Homo sapiens (subclone 3 e9 from P1 H55) DNA sequence	2e-007
		Sambucus nigra ribosome inactivating protein precursor	
1830	AF012899	mRNA, complete cds	2e-007
	www.acco	Sambucus nigra ribosome inactivating protein precursor	
***********	AF012899	mRNA, complete cds	9e-008
1832	L78776	Homo sapiens (subclone 2_a7 from P1 H49) DNA sequence	9e-008
	***************************************	Human ring zinc-finger protein (ZNF127-Xp) gene and 5'	
	U41315	flanking sequence.	9e-008
1834	X95586	H.sapiens MB1 gene	9e-008
1835	M33387	Human debrisoquine 4-hydroxylase (CYP2D8P) and	9e-008
1836	U09954	Human ribosomal protein L9 gene, 5' region and complete	8e-008
		H.sapiens gene for ventricular myosin light chain 2 > ::	
		gb L01652 HUMVMLC Human ventricular myosin light	man and
1837	Z15030	chain 2 gene, seven exons.	8e-008
		Sambucus nigra ribosome inactivating protein precursor	00-000
1838	AF012899	mRNA, complete cds	8e-008
		Sambucus nigra ribosome inactivating protein precursor	06-000
1839	AF012899	mRNA, complete cds	8e-008
1840	Z77974	H.sapiens flow-sorted chromosome 6 HindIII fragment,	8e-008
***************************************	***************************************	Sambucus nigra ribosome inactivating protein precursor	06-000
1841	AF012899	mRNA, complete cds	9- 009
	L81802	Homo sapiens (subclone 1_c12 from P1 H31) DNA	8e-008
	D87001	Human (lambda) DNA for immunoglobulin light chain	8e-008
		H. sapiens (D2S338) DNA segment containing (CA) repeat;	8e-008
1844	Z23971	clone AFM276zf5; single read	0 - 000
•••••	X89398	H.sapiens ung gene for uracil DNA-glycosylase	8e-008
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z68212	Phocine Herpesvirus 1 DNA (clone 4; 280 bp)	3e-008
······································	M85145	**	3e-008
-	M17919	Human tumor necrosis factor receptor, 3' flank.	3e-008
1040		Human DNA with homology to EBV IR3 repeat, clone Hu3.	3e-008
1840	M21339	Human non-histone chromosomal protein HMG-14 gene,	· ·
1077	19121339	complete cds.	3e-008
1850	Z69655	Human DNA sequence from cosmid L98A6, Huntington's	
1020		Disease Region, chromosome 4p16.3	3e-008
1051	202526	red photopigment gene {Alu repeat region, long intron 1}	***************************************
~~~~	S83526		3e-008
······································	M11809	Human (2'-5') oligo A synthetase E gene, exon 7 and flanks.	3e-008
1853	X94768	H.sapiens RP3 gene (XLRP gene 3)	3e-008
105		H.sapiens rod cG-PDE G gene for 3', 5'-cyclic nucleotide	***************************************
	X62025	phosphodiesterase	3e-008

SEQ ID	ACCESSION	DESCRIPTION	P VALU
		Homo sapiens aldehyde dehydrogenase 7 (ALDH7) mRNA	*******************************
	South States	>:: gb U10868 HSU10868 Human aldehyde dehydrogenase	***************************************
	5 NM_000694.1	ALDH7 mRNA, complete cds.	3e-008
	6 U22086	Ursus americanus clone G10H GT and ATTT microsatellite	3e-008
	7 AC001174	Homo sapiens (subclone 1_e2 from BAC H94) DNA	3e-008
	8 X69908	H.sapiens gene for mitochondrial ATP synthase c subunit	2e-008
**********************	9 X91233	H.sapiens IL15 gene	2e-008
NORTH CONTROL	M61835	Human lactase phlorizin hydrolase (LCT) gene, exon 2.	1e-008
-	M29324	Mouse L1Md-A13 repetitive sequence.	1e-008
1862	2 AF021806	Rattus norvegicus connexin 40 (GJA5) mRNA, complete cd	s 1e-008
		Rattus norvegicus mRNA for beta-alanine-pyruvate	
1863	AB002584	aminotransferase, complete cds	1e-008
	The state of the s	Human DNA sequence from cosmid L129H7, Huntington's	
1864	Z54147	Disease Region, chromosome 4p16.3 contains CpG island	9e-009
***************************************		Sambucus nigra ribosome inactivating protein precursor	70-009
1865	AF012899	mRNA, complete cds	9e-009
1866	Z54349	H.sapiens MN/CA9 GENE	8e-009
1867	M21487	Human muscle creatine kinase gene (CKMM), 5' flank.	3e-009
1868	U02993	Human cytochrome P450 (Cyp1A2) gene, 5' region.	3e-009
1869	Z78893	H.sapiens flow-sorted chromosome 6 HindIII fragment,	3e-009
	U89387	Human RNA polymerase II subunit hsRPB4 gene, complete	3e-009
	X57413	Mouse mRNA for transforming growth factor-beta2	3e-009
	Z94828	G.gallus microsatellite DNA (LEI0260	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
***************************************	D26067	Human mRNA for KIAA0033 gene, partial cds	3e-009
Contract contract and the contract cont	AB001914	Homo sapiens PACE4 gene, exon 23-25, complete cds	3e-009
***************************************		Human DNA sequence from cosmid U61F10, between	3e-009
1875	Z75894	markers DXS366 and DXS87 on chromosome X contains	2 - 000
***************************************	AC001443	Homo sapiens (subclone 2_f10 from BAC 2913	3e-009
	M96851	Human CpG island containing upstream sequence	3e-009
	D64108	Human mRNA for DMC1 homologue, complete cds	3e-009
••••••••••••••••••••••••••••••••••••••		{junction region} [human, KOPT-K1 cells, T-cell acute	3e-009
1879	S80861	lymphoblastic leukemia patient, Genomic, 895 nt	2 - 000
de transcer a company de la co	U79776	Mus musculus ajuba (Ajuba) mRNA, complete cds	3e-009
		thyrotropin-releasing hormone receptor [human, Genomic,	2e-009
1881	S75283	2312 nt, segment 2 of 2]	
Market Commission of the Commi	X14445	Human int-2 proto-oncogene	1e-009
···	J03764	Human, plasminogen activator inhibitor-1 gene, exons 2 to	1e-009
	L36911	Pig microsatellite DNA (CA repeat)	1e-009
		Homo sapiens serotonin transporter (hSERT) gene, promoter	1e-009
1885	U79746		
arena de la companya	X56668		9e-010
1000	2220000		9e-010
1887	AF012899	Sambucus nigra ribosome inactivating protein precursor	
1007	AT 012099	mRNA, complete cds	9e-010
1000	X77738	H.sapiens red cell anion exchanger (EPB3, AE1, Band 3)	
<del>~~~~~</del>		_	4e-010
arana arang ar	L02897 D45198	TT. Data C	3e-010
***************************************		TT	3e-010
1971	X04981	H. sapiens gene for lecithin-cholesterol acyltransferase	3e-010
1000	M14000	Human L1Heg repetitive element from the intergenic region	
and the second second second second second second	M14292	of the epsilon and G-gamma globin genes.	3e-010
1893	X14448	Human GLA gene for alpha-D-galactosidase A (EC	3e-010
1004	70.644.6	H.sapiens telomeric DNA sequence, clone 4QTEL025, read	
1894	Z96616	4QTELOO025.seq	3e-010

EQ ID	~\	DESCRIPTION	P VALU
189:	5[M12901	Human c-mos pseudogene with Alu repeat insertions, partial	
	Year Year	Human DNA sequence from cosmid L21F12B, Huntington's	
***************************************	5 Z68885	Disease Region, chromosome 4p16.3, contains EST	1e-010
1897	7¦L77036	Homo sapiens (subclone 5_d9 from P1 H19) DNA	1e-010
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	~*************************************
1898	Z58927	116g2, reverse read cpg116g2.rt1a	1e-010
*********	L77036	Homo sapiens (subclone 5_d9 from P1 H19) DNA	1e-010
1900	Z79007	H.sapiens flow-sorted chromosome 6 HindIII fragment,	1e-010
1901	Z75891	Human DNA sequence from cosmid F45C1 on chromosome	1e-010
1902	2 Z72930	S.cerevisiae chromosome VII reading frame ORF YGR145w	1e-010
1903	Z36111	S.cerevisiae chromosome II reading frame ORF YBR242w	5e-011
		Human DNA sequence from cosmid L129H7, Huntington's	***************************************
1904	Z54147	Disease Region, chromosome 4p16.3 contains CpG island	4e-011
1905	X87579	H.sapiens CD4 gene	4e-011
1906	U43604	Human unidentified mRNA, partial sequence.	4e-011
		Human clone A dehydroepiandrosterone sulfotransferase	
1907	U08024	(STD) mRNA, complete cds.	4e-011
1908	M27825	B.lactucae heat shock protein 70 (hsp70) gene, complete	4e-011
		H.sapiens genes for tumor necrosis factor (Tnfa) and	·
1909	Z15026	lymphotoxine (Tnfb)	3e-011
		Human DNA sequence from cosmid L206D7, Huntington's	
1910	Z67997	Disease Region, chromosome 4p16.3	3e-011
1911	AC001046	Homo sapiens (subclone 3_f2 from P1 H54) DNA sequence	2e-011
1912	Z84518	H.sapiens flow-sorted chromosome 6 HindIII fragment,	1e-011
1913	M59709	Human carcinoembryonic antigen (CEA) gene, exon 10.	1e-011
1914	L35670	Homo sapiens (subclone H8 10_g5 from P1 35 H5 C8) DNA	
		Sambucus nigra ribosome inactivating protein precursor	<del></del>
1915	AF012899	mRNA, complete cds	1e-011
		H.sapiens telomeric DNA sequence, clone 12PTEL055, read	***************************************
1916	Z96209	12PTELOO055.seq	1e-011
		Human Down Syndrome region of chromosome 21, genomic	***************************************
1917	U34052	sequence, clone A35A7-1A2.	1e-011
		H.sapiens telomeric DNA sequence, clone 21QTEL007,	
	Z96489	read 21QTELOO007.seq	1e-011
<del>~~</del> ~~~~~	AC001036	Homo sapiens (subclone 2_f7 from P1 H48) DNA sequence	1e-011
1920	L42098	Homo sapiens (subclone 5_c7 from P1 H22) DNA sequence.	6e-012
~~~~~	X93341	H.sapiens mitochondrial control region DNA	5e-012
1922	D26141	Human NF1 gene homologue	4e-012
		Human clotting factor VIII gene, intron 20 and exon 21,	
**************	U80228	partial sequence	4e-012
1924	U16812	Human Bak-2 gene, complete cds.	4e-012
		Human DNA sequence from cosmid cN85E10 on	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z68758	chromosome 22q11.2-qter	4e-012
***************************************	AC001449		4e-012
1927	D50561		4e-012
		H.sapiens telomeric DNA sequence, clone 18PTEL033, read	
1928	Z96387	10DELL COOC	1e-012
		Homo sapiens 16S ribosomal RNA, mitochondrial gene,	
1929	AF004338	i . •	1e-012
		Human transposon-like element (THE) p2 solo LTR with	
1930	M15360	· · · · · · · · · · · · · · · · · · ·	1e-012
1931	L81577	\$	1e-012
1020	U56979		1e-012

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
***************************************	U14550	Human sialyltransferase SThM (sthm) mRNA, complete cds.	1e-012
1934	M31061	Human ornithine decarboxylase gene, complete cds.	1e-012
		Rattus norvegicus Fisher 344 pre-sialomucin complex	V V (100000)
1935	U06752	(pSMC) mRNA, repeat sequences 10-14, partial cds.	1e-012
		Rattus norvegicus Fisher 344 pre-sialomucin complex	
MANAGEMENT AND	U06752	(pSMC) mRNA, repeat sequences 10-14, partial cds.	1e-012
	L29096	Homo sapiens oriP binding protein (OBP-2) mRNA,	1e-012
1938	U50156	Human DNA segment containing CA repeat at locus	5e-013
		H.sapiens RRM1 gene for ribonucleoside diphosphate	-
***************************************	X65708	reductase M1 subunit	5e-013
1940	M61107	Human p22-phox (CYBA) gene, exons 3 and 4.	5e-013
1941	AC001502	Homo sapiens (subclone 2_c7 from P1 H43) DNA sequence	4e-013
		H.sapiens gene for casein kinase II alpha subunit > subunit	Control of the contro
1942	X69951	alpha [human, Genomic, 18862 nt]	4e-013
1943	AC002252	Homo sapiens (subclone 1 g7 from BAC H76) DNA	4e-013
	······································	Caenorhabditis elegans cosmid C45B11, complete sequence	•
1944	Z74029	[Caenorhabditis elegans]	4e-013
1945	U90544	Human sodium phosphate transporter (NPT3) mRNA,	4e-013
1946	L77032	Homo sapiens (subclone 3_e5 from P1 H16) DNA sequence.	
····	X55367	Human alpha-satellite DNA from clone pTRA-2	2e-013
	U40369	Human spermidine/spermine N1-acetyltransferase	2e-013
	Z25749	H.sapiens gene for ribosomal protein S7	2e-013
***************************************	M96838	Human proteinase 3 gene, exons 2 and 3.	2e-013
	Z73116	S.cerevisiae chromosome XII reading frame ORF YLL011w	
ON THE RESIDENCE OF THE PARTY O	U90544	Human sodium phosphate transporter (NPT3) mRNA,	2e-013
***************************************	U90544	Human sodium phosphate transporter (NPT3) mRNA,	2e-013
~~~~~~	AC001016	Homo sapiens (subclone 2 f8 from P1 H43) DNA sequence	1e-013
		Homo sapiens gene for thymidylate synthase, exons 1, 2, 3,	
1955	D00596	4, 5, 6, 7, complete cds	1e-013
	X80240	H.sapiens endogenous retrovirus HERV-KC4 DNA	1e-013
····	M86181	Human prosaposin (PSAP) gene.	5e-014
·····		Human mitochondrial genes for several tRNAs (Phe, Val,	
1958	V00710	Leu) and 12S and 16S ribosomal RNAs	5e-014
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	30 014
1959	Z62151		5e-014
		Homo sapiens interleukin 12B (natural killer cell stimulatory	30-01-4
		factor 2, cytotoxic lymphocyte maturation factor 2, p40)	
		(IL12B) mRNA > :: gb M65290 HUMNKSFP40 Human	
1960	NM 002187.1		5e-014
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M18680	~_~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	5e-014
		Human endogenous retrovirus-like sequence (LTR ERS-P1-	30-01-4
1962	X06274		5e-014
	2 2 0 0 2 7 1	Human immunodeficiency virus type 2 proviral DNA,	JC-014
1963	D00835	;	5e-014
		Human Alu repetitive sequence located near the insulin gene	JC-014
and the same of th		>:: gb J00268 HUMINS02 Human insulin gene: repetitive	
1964	V00581		50.014
1704	¥ 00301	Human DNA sequence from cosmid A1 on chromosome 6	5e-014
1065	Z95437		E - 014
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	AB001051	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	5e-014
·····		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	5e-014
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	AB001051		5e-014
<del>เพราะสาราสาราสาราสาราสาราสาราสาราสาราสาราส</del>	M59709	The state of the s	2e-014
1969	X91233	H.sapiens IL15 gene	2e-014

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
VANAGE	X12718	Human Retrovirus mRNA for LTR (clone cPB-3)	
1970	A12/10	Human interferon genes LeIF-L and LeIF-J and pseudogene	2e-014
1971	V00531	LeIF-M with intergenic regions. These genes are located on	1 - 014
1771	V 00331	Human L1Heg repetitive element from the intergenic region	1e-014
1972	M14292	of the epsilon and G-gamma globin genes.	1.0014
1712	14114272	Human UbA52 adrenal mRNA for ubiquitin-52 amino acid	1e-014
1973	X56998	fusion protein	1e-014
	S74906	E1 beta=pyruvate dehydrogenase beta {promoter}	8e-015
	M18680	Homo sapiens 5S rRNA pseudogene.	6e-015
	L49046	Homo sapiens (subclone 2 h3 from P1 H25) DNA sequence	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M18680	Homo sapiens 5S rRNA pseudogene.	6e-015
	11110000	Homo sapiens elastin gene, exons 5-27 and alternatively	00-013
1978	U93037	spliced products, partial cds	6e-015
warman and a second	M76741	Homo sapiens biliary glycoprotein (BGP) gene, partial cds.	6e-015
	14107-11	Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	06-013
1980	U72787	complete sequence [Homo sapiens]	6e-015
#49444995717114.040404040400000	AB000931	Homo sapiens FUT2 gene, intron 1, complete sequence	6e-015
~~~~~~~	M83137	Human scaffold-attached region (SAR) DNA.	6e-015
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M18680	Homo sapiens 5S rRNA pseudogene.	5e-015
	11110000	H.sapiens CpG island DNA genomic Mse1 fragment, clone	36-013
1984	Z63454	84d2, reverse read cpg84d2.rt1a	5e-015
1701	203737	H.sapiens CpG island DNA genomic Mse1 fragment, clone	26-013
1985	Z63454	84d2, reverse read cpg84d2.rt1a	5e-015
	X97489	H.sapiens PIT1/GHF1 gene silencer region	5e-015
	X55367	Human alpha-satellite DNA from clone pTRA-2	2e-015
and the second s	Z22795	H.sapiens microsatellite repeat	2e-015
***************************************	D38112	Human mitochondrial DNA, complete sequence	2e-015
		H.sapiens gene for casein kinase II alpha subunit subunit	20-013
1990	X69951	alpha [human, Genomic, 18862 nt]	2e-015
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	M59250	Homo sapiens cytochrome c oxidase subunit Vb	2e-015
······································	X15965	Rabbit DNA for L10c5 repeat	2e-015
-	X70052	S.cerevisiae sof1 gene	2e-015
contraction in the contraction of the	M86667	H.sapiens NAP (nucleosome assembly protein) mRNA,	1e-015
·····	X78212	H.sapiens diamine oxidase gene	7e-016
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U21928	Human fructose-1,6-biphosphatase (FBP1) gene, exon 4	7e-016
**************************************	AB001051	Dugesia japonica mRNA for ADP-ribosylation factor,	7e-016
ananamanan sanas	AC002181	Homo sapiens (subclone 2 a12 from BAC H111) DNA	6e-016
		H.sapiens DNA for X-linked dinucleotide repeat	00 010
1999	X91413	polimorphism (clone YCA3CL149)	6e-016
		Human ribosomal RNA upstream binding transcription	
2000	U65487	factor (UBTF) gene, partial cds	6e-016
***************************************		Human DNA sequence from cosmid 91K3, Huntington's	
2001	Z69920	Disease Region, chromosome 4p16.3	6e-016
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		Homo sapiens c33.28 unnamed HERV-H protein mRNA,	6e-016
2003	CONTRACTOR OF THE PROPERTY OF	Human colin carcinoma laminin-binding protein mRNA,	3e-016
		Human DNA sequence from cosmid L21F12B, Huntington's	
2004	Z68885	<u>.                                    </u>	2e-016
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	***************************************	Homo sapiens apoptosis inhibitor survivin gene, complete	2e-016
· COLONIA ARABAMAN AND AND AND AND AND AND AND AND AND A	M23442	Human interleukin 4 (IL-4) gene, complete cds.	2e-016
		Human thymopoietin (TMPO) gene, exons 4 and 5, and	
2007	U18270	complete cds for thymopoietin alpha	2e-016
		Homo sapiens DNA from chromosome 19-cosmid f19399	
2008	AD000833		2e-016
		· · · · · · · · · · · · · · · · · · ·	

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
		Homo sapiens DNA from chromosome 19-cosmid f19399	
2009	AD000833	(~17 kb EcoRI restriction fragment)	2e-016
2007	215000055	Drosophila melanogaster SURF-4 gene and gene encoding	20-010
2010	Y14823	seryl-tRNA synthetase	2e-016
2010		'Homo sapiens intestinal mucin (MUC2) gene, promoter	20 010
2011	U67167	region and partial cds	8e-017
		H.sapiens telomeric DNA sequence, clone 10QTEL040,	00-017
2012	Z96177	read 10QTELOO040.seq	7e-017
2012		Homo sapiens cytosolic phagocyte oxidase protein	70-017
2013	AF003533	(p47phox) gene, promoter region and partial cds	7e-017
2013	A1003333	Homo sapiens glucose-6-phosphatase, catalytic glucose-6-	76-017
		phosphatase mRNA, complete cds. >:: gb[I15157]I15157	
2014	NM 000151.1	Sequence 1 from patent US 5460942	7e-017
NATURAL DESCRIPTION DE LA CONTRACTION DE LA CONT	X94912	H.sapiens Pr22 gene	7e-017
2013	A)4912	Human junction sequence from chimeric/rearranged YAC	76-017
2016	U10580	yRM2252, chromosome 11p14.	7e-017
~~~~~~~~	M22970	Human pancreatic phospholipase A-2 (PLA-2) gene, exons 1	7e-017
	X02152	Human mRNA for lactate dehydrogenase-A (LDH-A, EC	6e-017
2016	AU2132	Human DNA sequence from cosmid L118D5, Huntington's	06-017
2010	Z68755	Disease Region, chromosome 4p16.3	6- 017
2019	208/33	Human microsatellite DNA repeat region DNA. > ::	6e-017
2020	N#64904	· · · · · · · · · · · · · · · · · · ·	2- 017
	M64804 Z92910	gb  I31115  I31115 Sequence 27 from patent US 5582979	3e-017
ann an	X74984	H. sapiens HFE (HLA-H) gene	3e-017
	U05333	H.sapiens 5' flanking region of CD14 gene	2e-017
	U48485	Mus musculus co-chaperonin 'cofactor A' mRNA, complete Human skeletal muscle ryanodine receptor gene	2e-017
	X97869		2e-017 2e-017
	X17579	H.sapiens gene encoding La autoantigen Human specific HS5 DNA	2e-017 2e-017
university of the second second	U36445	Bos taurus calcium-activated chloride channel mRNA,	2e-017
recommendation and a second and a	L06845	Human cysteinyl-tRNA synthetase mRNA, partial cds.	2e-017
	X93334	H.sapiens mitochondrial DNA, complete genome	8e-018
en moneron anno anno anno anno anno anno anno a	X62996	H.sapiens mitochondrial genome (consensus sequence)	
NAMES OF THE PARTY	M98479	Human transglutaminase mRNA, 3' untranslated region.	8e-018 8e-018
2031	101704/7	Homo sapiens (clone ISW11-1) non-gastric H,K-ATPase	86-019
2032	L42568	(ATP1AL1) gene, exons 15-17	8e-018
	S52659		<u> </u>
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	U66707	lactoferrin {5' region, promoter} [human, placenta, Rattus norvegicus densin-180 mRNA, complete cds	8e-018
<del>arraman arraman arram</del>	X76683	Plasmid vector pHM2 betalactamase gene	8e-018
announce and a second	U66707	Rattus norvegicus densin-180 mRNA, complete cds	3e-018 3e-018
	D17201	Human HepG2 3' region MboI cDNA, clone hmd3d04m3	3e-018
erconoucumous xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx	Z24281	H. sapiens (D12S355) DNA segment containing	2e-018
	AC001443	Homo sapiens (Subclone 2 fl0 from BAC 2913	2e-018 2e-018
~~~~~~	U73522	Homo sapiens AMSH mRNA, complete cds	2e-018
••••••••••••••••••••••••••••••••••	L77040	Homo sapiens (subclone 8 c11 from P1 H22) DNA	9e-019
	**************************************		C
2042	X93496	H.sapiens TRAP gene, 5' flanking region Human gene for alpha-1-microglobulin-bikunin, exons 1-5	9e-019
2042	V5/1816		0- 010
***************************************	X54816	(encoding alpha-1-microglobulin, N-terminus.)	9e-019
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	L35240	Human enigma gene, complete cds	8e-019
2043	Z46940	H.sapiens PRM1 gene, PRM2 gene and TNP2 gene	8e-019
2046	11/0001 1	Human poly(A)-binding protein (PABP) processed	0.010
2046	U60801.1	pseudogene2, complete cds	8e-019
•••••••••••••••••••••••••••••••••••••••		Homo sapiens nucleotide binding protein 1 Human	£

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
	X76683	Plasmid vector pHM2 betalactamase gene	3e-019
2049	U40369	Human spermidine/spermine N1-acetyltransferase	3e-019
2050	D13624	Human Wilms tumor gene encoding WT1 protein, exon 6	3e-019
2051	D13624	Human Wilms tumor gene encoding WT1 protein, exon 6	3e-019
2052	D86566	Human DNA for NOTCH4, partial cds	3e-019
		Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	
2053	U72787	complete sequence [Homo sapiens]	3e-019
2054	U86759	Human netrin-2 like protein (NTN21) mRNA, complete cds	3e-019
2055	X78454	X.laevis AB21 mRNA for RPD3 homologue	3e-019
2056	NM 000969.1	Homo sapiens ribosomal protein L5 (RPL5) mRNA	1e-019
2057	U49869	Human ubiquitin gene, complete cds	1e-019
2058	X78901	H.sapiens (lambda63) DNA of apolipoprotein cluster	9e-020
nicompression in an exercise or a	in series are transported and acceptance and accept	Human putative tumor suppressor (MXI1) gene, exons 4, 5,	\$~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
2059	U32515	and 6, and complete cds	9e-020
	D87717	Human mRNA for KIAA0013 gene, complete cds	5e-020
		Human DNA sequence from cosmid cN85E10 on	-
2061	Z68758	chromosome 22q11.2-qter	3e-020
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Z30584	R.norvegicus (wistar) mRNA for ZG-16p	3e-020
2002	230304	Human DNA sequence from cosmid cN85E10 on	JC-020
2063	Z68758	chromosome 22q11.2-qter	3e-020
	M58600	Human heparin cofactor II (HCF2) gene, exons 1 through 5.	3e-020
2004	14136000	Homo sapiens elastin gene, exons 5-27 and alternatively	36-020
2065	1102027		2- 020
***************************************	U93037	spliced products, partial cds	3e-020
**********************	L38951	Homo sapiens importin beta subunit mRNA, complete cds	2e-020
2007	D26141		1e-020
20.60	N410706	Orangutan beta- and delta-globin gene intergenic region with	3
	M18796	2 Alu repeats.	1e-020
0.40.46.46.00.40.40.40.40.40.40.40.40.40.	Z47046	Human cosmid QLL2C9 from Xq28	1e-020
-	AC001443	Homo sapiens (subclone 2_f10 from BAC 2913	1e-020
20/1	X15965	Rabbit DNA for L1Oc5 repeat	1e-020
	*****	Human phosphatidylinositol 3-kinase catalytic subunit	
THE RESIDENCE PROPERTY OF THE PERSON ASSESSMENT ASSESSM	U86453	p110delta mRNA, complete cds	6e-021
	M90058	Human serglycin gene, exons 1,2, and 3.	4e-021
***	L81932	Homo sapiens (subclone 9_h2 from P1 H21) DNA sequence	
************	X68258	Bicistronic transcription units (pSBC-2)	3e-021
2076	L78777	ифинальностического <del>на</del> ничистойни финальностического <del>и н</del> а съставани от честически и на на нестически били вы начина	3e-021
	J01415	Human mitochondrion, complete genome	1e-021
	L43411	Homo sapiens (subclone 5_g5 from P1 H25) DNA sequence.	1e-021
2079	NM_000967.1	<sup>1</sup> Homo sapiens ribosomal protein L3 (RPL3) mRNA protein	1e-021
		Human DNA sequence from cosmid U65A4, between	
2080	Z81014	markers DXS366 and DXS87 on chromosome X *	1e-021
		Homo sapiens elastin gene, exons 5-27 and alternatively	
2081	U93037	spliced products, partial cds	1e-021
2082	X78454	X.laevis AB21 mRNA for RPD3 homologue	1e-021
2083	X82825	A.thaliana PRL1 mRNA	1e-021
**********************	J01415	Human mitochondrion, complete genome	4e-022
······································		Human DNA sequence from cosmid L21F12B, Huntington's	***************************************
2085	Z68885	Disease Region, chromosome 4p16.3, contains EST	4e-022
****************************	L81690	พริกาศาสตราชาวาราสาธารรรมที่ พระสาธารรมสาธารรมสาธารรมสาธารรมสาธารรมสาธารรมสาธารรมสาธารรมสาธารรมสาธารรมสาธารรมส	4e-022
	Z47046	Human cosmid QLL2C9 from Xq28	4e-022
		Human DNA sequence from cosmid 92M18, BRCA2 gene	
2088	Z73360	region chromosome 13q12-13	1e-022

SEQ ID	ACCESSION	DESCRIPTION	P VALUE
	ana	H.sapiens telomeric DNA sequence, clone 12PTEL057, read	
2090	Z96210	12PTELOO057.seq	5e-023
	Z50751	H.sapiens mRNA for B4B	4e-023
	U14567	***ALU WARNING: Human Alu-J subfamily consensus	4e-023
	L35657	Homo sapiens (subclone H8 5_a10 from P1 35 H5 C8) DNA	4e-023
***************	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Human DNA sequence from fosmid F62D4 on chromosome	
		22q12-qter > :: emb Z81316 HSF62D4A Human DNA	
2094	Z81315	1	4e-023
CALLED AND AND AND AND AND AND AND AND AND AN	U93237	Human menin (MEN1) gene, complete cds	1e-023
***	M16660	Human 90-kDa heat-shock protein gene, cDNA, complete	1e-023
	Z93943	Human DNA sequence from cosmid U235H3 on	6e-024
	M84334	Macacca mulatta hnRNP A1-gamma isoform mRNA,	5e-024
2098	14104334	Human DNA sequence from cosmid 92M18, BRCA2 gene	30-024
2000	772260	•	5e-024
	Z73360	region chromosome 13q12-13	5e-024
	L32754	Homo sapiens Ig-associated signalling molecule	5e-024
2101	AC002252	Homo sapiens (subclone 1_g7 from BAC H76) DNA	3e-024
		Homo sapiens ribosomal protein L21 (RPL21) mRNA > ::	
		gb U25789 HSU25789 Human ribosomal protein L21	
2102	NM_000982.1	mRNA, complete cds.	5e-024
		Homo sapiens cytochrome b-245, beta polypeptide encoding	
0		mitochondrial protein, mRNA > :: emb X04011 HSXCGD	
		Human mRNA of X-CGD gene involved in chronic	9
2103	NM_000397.1	granulomatous disease located on chromosome X	5e-024
2104	L19086	Human LINE1 (L1.3) repetitive element DNA sequence.	4e-024
2105	U88531	Bos taurus phosphatidylinositol 4-kinase mRNA, complete	2e-024
2106	D38112	Human mitochondrial DNA, complete sequence	2e-024
	} !	Homo sapiens SET translocation (myeloid leukemia-	
2107	NM 003011.1	associated) (SET) mRNA > :: gb M93651 HUMSET Human	2e-024
2108	M98512	Human NFG genomic fragment.	2e-024
2109	NM 001019.1	Homo sapiens ribosomal protein S15a for ribosomal protein	2e-024
·····	X70991	H.sapiens MADER mRNA	2e-024
	X14445	Human int-2 proto-oncogene	2e-024
	AC001174	Homo sapiens (subclone 1 e2 from BAC H94) DNA	2e-024
	D86566	Human DNA for NOTCH4, partial cds	2e-024
	Z78885	H.sapiens flow-sorted chromosome 6 HindIII fragment,	1e-024
	X82272	Human endogenous retrovirus env mRNA	1e-024
***********		H.sapiens telomeric DNA sequence, clone 10QTEL017,	
2116	Z96167	read 10QTELOO017.seq	6e-025
	L38951	Homo sapiens importin beta subunit mRNA, complete cds	6e-025
	X53575	Yeast RPL7 gene for ribosmal protein L7	6e-025
CONTRACTOR OF THE STATE OF THE	L77040	Homo sapiens (subclone 8 c11 from P1 H22) DNA	5e-025
2117		H. sapiens (D2S336) DNA segment containing (CA) repeat;	
2120	Z23957	clone AFM275yf5; single read	5e-025
***********************	}	Human int-2 proto-oncogene	5e-025
	X14445	Homo sapiens interleukin 9 receptor IL9R pseudogene,	2e-025
2122	L39062	Human DNA sequence from phage LAW2 from a contig	ZC-UZJ
	4		
2422	70.4702	from the tip of the short arm of chromosome 16, spanning	2- 025
************	Z84723	2Mb of 16p13.3 Contains Interleukin 9 receptor pseudogene	
2124	AC001174	Homo sapiens (subclone 1_e2 from BAC H94) DNA	2e-025
		Homo sapiens epithelial membrane protein 1 Progression	
	NM_001423.1	Associated Protein	2e-025
2126	X14445	Human int-2 proto-oncogene	2e-025

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
		Human L1 repetitive sequence with a region homologous to	STATE OF THE OWNERS AND THE STATE OF THE STA
2127	M68841	a mouse ORF.	2e-025
	X97489	H.sapiens PIT1/GHF1 gene silencer region	2e-025
	X73501	H.sapiens gene for cytokeratin 20	1e-025
	U48363	Mus musculus transcriptional activator alpha-NAC	1e-025
	X01037	Human 7SL RNA sequence	7e-026
	M22485	Human ring chromosome 21 [r(21)] breakpoint DNA.	2e-026
	K03429	Ape (chimpanzee) 28S ribosomal RNA gene.	2e-026
4177	NUJTZJ	Homo sapiens cellular apoptosis susceptibility protein	120 020
2134	AF053644	(CSE1) gene, exon 2	9e-027
	L06900	Human dystrophin gene, intron 1 containing pseudo exon.	7e-027
2133	L00900	Human thymopoietin (TMPO) gene, partial exon 6,	70-027
2126	T.1.0071		7- 027
	U18271	complete exon 7, partial exon 8, and partial cds for	7e-027
	X05323	Human MRC OX-2 gene signal sequence	7e-027
	M99065	Rat core histone (MacroH2A.1) mRNA, complete cds.	7e-027
augustasian sanan anatan salah salah salah s	X01037	Human 7SL RNA sequence	6e-027
	M22485	Human ring chromosome 21 [r(21)] breakpoint DNA.	2e-027
2141	D50494	Mouse mRNA for murine RCK, complete cds	2e-027
	A.A.V. TAN	Human L1Heg repetitive element from the intergenic region	Action of the second
2142	M14292	of the epsilon and G-gamma globin genes.	2e-027
		Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	***************************************
2143	U72787	complete sequence [Homo sapiens]	7e-028
		Homo sapiens ribosomal protein S27 gb L19739 HUMMPSI	
	novel control	Homo sapiens metallopanstimulin (MPS1) mRNA, complete	oosia oo
2144	NM 001030.1	cds. > :: gb I60224 I60224 Sequence 1 from patent US	7e-028
	AC002186	Homo sapiens (subclone 1 fl2 from P1 H115) DNA	7e-028
	M22485	Human ring chromosome 21 [r(21)] breakpoint DNA.	3e-028
***************************************	<u>.</u>	Human DNA sequence from cosmid A1 on chromosome 6	
2147	Z95437	contains ESTs. HERV like retroviral sequence	2e-028
		Homo sapiens ribosomal protein S23 (RPS23) mRNA > ::	
	300	dbj D14530 HUMRSPT Human homolog of yeast ribosomal	NO TO THE PERSON OF THE PERSON
2148	NM 001025.1	protein S28, complete cds	2e-028
2170	14141 001025.1	Human DNA sequence from cosmid A1 on chromosome 6	
2140	Z95437	contains ESTs. HERV like retroviral sequence	2e-028
2147	Z93431	H.sapiens CpG island DNA genomic Mse1 fragment, clone	20-020
2150	Z62151	64c7, forward read cpg64c7.ft1a	1e-028
2130	202131	H.sapiens CpG island DNA genomic Mse1 fragment, clone	16-026
2161	760151	64c7, forward read cpg64c7.ft1a	8e-029
AND	Z62151		Egypte, to the telescope and the contract of t
2152	X62996	H. sapiens mitochondrial genome (consensus sequence)	8e-029
0150	760151	H.sapiens CpG island DNA genomic Mse1 fragment, clone	0. 000
	Z62151	64c7, forward read cpg64c7.ft1a	8e-029
	AC001518	Homo sapiens (subclone 2_b4 from P1 H49) DNA sequence	8e-029
	X56932	H.sapiens mRNA for 23 kD highly basic protein	8e-029
2156	U93563	Human L1 element L1.6 putative p150 gene, complete cds	8e-029
	***************************************	H.sapiens telomeric DNA sequence, clone 13QTEL058,	*
2157	Z96282	read 13QTELOO058.seq	8e-029
	The state of the s	Homo sapiens retinoblastoma-binding protein 1 (RBBP1)	
	Acceptance of the Control of the Con	mRNA > :: gb S66427 S66427 RBP1=retinoblastoma	5
2158	NM_002892.1	binding protein 1 [human, Nalm-6 pre-B cell leukemia,	7e-029
		Human endogenous retrovirus DNA downstream of 5' LTR,	***

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
	nassan indragripanyanidan indragripanian indragripanian indragripanian indragripanian indragripanian indragrip	Human DNA sequence from intron 22 of the factor VIII	200000000000000000000000000000000000000
		gene, Xq28. Contains the end of a 9.5kb repeated region,	
2160	X86012	int22h-1, involved in many cases of haemophilia	3e-029
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	miczii i, myorved m many dusts si	3e-029
	L81840	H.sapiens gene for alpha-2 macroglobulin, exon 1	3e-029
	Z11711	11.3apiciis gene for dipita 2 marti-8-1-1	3e-029
	X61453	Homo sapiens WD repeat domain 2 (WDR2) mRNA >	1e-029
	NM_003389.1	Homo sapiens wD repeat domain 2 (wDR2) likiNA	9e-030
2165	L81840	TIOTIO Supicità (Subciente 1 10 11 0111 1 1 1 1 1 1 1 1 1 1 1 1 1	9 <b>C</b> -030
		Homo sapiens chromosome 21 DNA fragment with Alu, L1	8- 020
2166	L05173		8e-030
		Human endogenous retrovirus DNA downstream of 5' LTR,	2 020
2167	M12855	clone HERV-K22.	3e-030
	(CO) (CO) (CO) (CO) (CO) (CO) (CO) (CO)	Human endogenous retrovirus DNA downstream of 5' LTR,	
2168	M12855	clone HERV-K22.	3e-030
2169	J05211	Human desmoplakin mRNA, 3' end.	3e-030
***************************************	<u></u>	H.sapiens telomeric DNA sequence, clone 10QTEL017,	
2170	Z96167	read 10OTELOO017.seq	3e-030
		Human endogenous retrovirus DNA downstream of 5' LTR,	
2171	M12855	clone HERV-K22.	3e-030
	AC002186	Homo sapiens (subclone 1_f12 from P1 H115) DNA	3e-030
ALL DESCRIPTIONS OF THE PARTY O	L35657	Homo sapiens (subclone H8 5_a10 from P1 35 H5 C8) DNA	3e-030
	Survey and the survey of the s	Homo sapiens (subclone H8 5_a10 from P1 35 H5 C8) DNA	3e-030
	L35657	Homo sapiens transcription factor SL1 mRNA, partial cds.	3e-030
	L39061	Homo sapiens Pig8 (PIG8) mRNA, complete cds	1e-030
Maria de la companione	AF010313	Homo sapiens (subclone 2_f10 from BAC 2913	1e-030
2177	AC001443	Homo sapiens (subcione 2 110 from BAC 2915	10-030
	establishment.	H.sapiens CpG island DNA genomic Mse1 fragment, clone	3e-031
	Z62151	64c7, forward read cpg64c7.ft1a	á
	AF010312	Homo sapiens Pig7 (PIG7) mRNA, complete cds	1e-031
	U77085	Human epithelial membrane protein (CL-20) mRNA,	1e-031
2181	D29805	Human mRNA for beta-1,4-galactosyltransferase, complete	1e-031
		Human DNA sequence from cosmid A1 on chromosome 6	
2182	Z95437	contains ESTs. HERV like retroviral sequence	1e-031
2183	X62996	H.sapiens mitochondrial genome (consensus sequence)	1e-031
		Lactococcus lactis cremoris glucose inhibited division	
2184	U80409	protein homolog GidA (gidA) gene, partial cds	4e-032
	X00525	Mouse 28S ribosomal RNA	1e-032
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		Human endogenous retrovirus clone c5.11, HERV-H	***************************************
2186	U35032	multiply spliced subgenomic leader, protease and integrase	1e-032
		Human DNA sequence from cosmid U61F10, between	
2187	7 Z75894	markers DXS366 and DXS87 on chromosome X contains	1e-032
	U22055	Human 100 kDa coactivator mRNA, complete cds.	1e-032
	M98509	Human NFB genomic fragment.	5e-033
	D38112	Human mitochondrial DNA, complete sequence	2e-033
2190	שנים על 112	Human DNA sequence from cosmid L96F8, Huntington's	<del></del>
	000	Disease Region, chromosome 4p16.3 contains EST and	orene water
		cDNA > :: emb Z69365 HSL96F8A Human DNA sequence	
	****		and the second
	Water Company	from cosmid L96F8, Huntington's Disease Region,	50.024
219	1 Z69364	chromosome 4p16.3 contains EST and cDNA	5e-034
	Steener answers	Human apurinic/apyrimidinic endonuclease (HAP1) gene, 5'	
	2 U59695	upstream region	5e-034
2193	3 L35657	Homo sapiens (subclone H8 5_a10 from P1 35 H5 C8) DNA	2e-034
2194	4 L78778	Homo sapiens (subclone 2_e10 from P1 H49) DNA	2e-034
The same of the sa	5 V00662	H.sapiens mitochondrial genome	5e-035

SEQ ID	ACCESSION	DESCRIPTION	P VALU
2196	M30629	Human pregnancy-specific glycoprotein beta-1	5e-035
-		Homo sapiens signal recognition particle 14kD (homologous	
2197	NM 003134.1	Alu RNA-binding protein) (SRP14) mRNA > recognition	5e-035
	U93572	Human L1 element L1.25 p40 and putative p150 genes,	2e-035
	a annuarium municum en electrica de la companie de	H.sapiens telomeric DNA sequence, clone 10QTEL038,	
2199	Z96176	read 10QTELOO038.seq	6e-036
·····	······································	Human mitochondrial genes for several tRNAs (Phe, Val,	
2200	V00710	Leu) and 12S and 16S ribosomal RNAs	6e-036
	Z78715	H.sapiens flow-sorted chromosome 6 HindIII fragment,	5e-036
	U93574	Human L1 element L1.39 p40 and putative p150 genes,	2e-036
	S63912	D10S102=FBRNP [human, fetal brain, mRNA, 3043 nt]	1e-036
announce and the second	AB011137	Homo sapiens mRNA for KIAA0565 protein, complete cds	1e-036
2204	ADVIII3/	H.sapiens CpG island DNA genomic Mse1 fragment, clone	
2205	Z55370	37a7, forward read cpg37a7.ft1a	7e-037
	L36720	Homo sapiens bystin mRNA, complete cds	7e-037
2200	L30/20	HTS1=HeLa tumor suppressor gene [human, revertant clone	
2207	045026	F2, mRNA Partial, 2687 nt]	7e-037
~~~~~~~~	S45936	Human thrombin receptor (F2R) gene, 5' region and partial	6e-037
and the second	U36755	Human infolioni receptor (12K) gene, 5 region and partial	3e-037
	ZZUJ:DJUJ-T		2e-037
	AB001325	Human AQP3 gene for aquaporine 3 (water channel), partail Homo sapiens (subclone H8 5_a10 from P1 35 H5 C8) DNA	
	L35657	Homo sapiens (subcione H8 5 allo from P1 55 H5 C8) DIVA	2e-037
	U36445	Bos taurus calcium-activated chloride channel mRNA,	7e-038
	X66292	P.pygmaeus (OX3910-11) alphoid repetitive DNA	
2214	D38112	Human mitochondrial DNA, complete sequence	6e-038
		H. sapiens (D12S80) DNA segment containing (CA) repeat;	C= 029
	Z16571	clone AFM102xd6; single read	6e-038
	D50561	Human DNA, replication enhancing element (REE1)	3e-038
2217	X66292	P.pygmaeus (OX3910-11) alphoid repetitive DNA	8e-039
		Human DNA sequence from cosmid 92M18, BRCA2 gene	7 020
	Z73360	region chromosome 13q12-13	7e-039
2219	M25718	Human rDNA and 4 Alu repeats.	4e-039
		Homo sapiens cytochrome b-245, beta polypeptide encoding	
	AVA-	mitochondrial protein, mRNA > :: emb X04011 HSXCGD	
	were the second	Human mRNA of X-CGD gene involved in chronic	
2220	NM 000397.1	granulomatous disease located on chromosome X	3e-039
2221	M13073	Human metallothionein I processed pseudogene, complete	2e-039
······································	Z78715	H.sapiens flow-sorted chromosome 6 HindIII fragment,	3e-040
	<u>.</u>	Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	
2223	U72787	complete sequence [Homo sapiens]	3e-040
	Z12962	H.sapiens mRNA for homologue to yeast ribosomal protein	3e-040
regression and regression (AC NO SEC SEC AND ARCHITECTURE)	Z78715	H.sapiens flow-sorted chromosome 6 HindIII fragment,	9e-041
	AC001443	Homo sapiens (subclone 2_f10 from BAC 2913	4e-041
		Human DNA sequence from cosmid 92M18, BRCA2 gene	<u> </u>
2227	Z73360	region chromosome 13q12-13	3e-041
	S72304	rah=ras-related homolog [mice, HT4 neural cell line,	3e-041
	AC001443	Homo sapiens (subclone 2 fl0 from BAC 2913	1e-041
	U49974	Human mariner2 transposable element, complete consensus	1e-041
	danaanaan ahaa ka k	Mouse myocyte nuclear factor (MNF) mRNA, complete cds.	
NORTH TO SERVICE AND THE PROPERTY OF THE PROPE	L26507	Human R kappa B mRNA, complete cds.	1e-042
	U08191	Homo sapiens RGH2 gene, retrovirus-like element	1e-042
	D11078 U83908	Human nuclear antigen H731 mRNA, complete cds	1e-042
			⇒ ↓ し⁻ひてム

Table 2A Nearest	Neighbor	(BlastN vs.	GenBank)

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
	0. 3. 45 Marie - 10. 10. 10. 10. 10. 10. 10. 10. 10. 10.	Homo sapiens protein kinase C substrate 80K-H (PRKCSH)	i
		mRNA > :: gb J03075 HUMG19P1A Human 80K-H protein	*
2236	NM 002743.1	(kinase C substrate) mRNA, complete cds.	1e-042
*****************************	L78777		4e-043
	BANTON MATERIA MATERIANA PARAMETER METALON METALON PARAMETER A ASSESSE AND ASSESSED AS A METALON ASSESSED AS A	H. sapiens CpG island DNA genomic Mse1 fragment, clone	
2238	Z59382	152b10, reverse read cpg152b10.rt1a	1e-043
****	L12469	Gallus gallus (max) gene, complete cds.	1e-043
***************************************		Homo sapiens cosmid clone U197H5 from Xp22.1-22.2,	
2240	U72789	complete sequence [Homo sapiens]	5e-044
······	·	Oryctolagus cuniculus peroxisomal Ca-dependent solute	1
2241	AF004161	carrier mRNA, complete cds	5e-044
	M22485	Human ring chromosome 21 [r(21)] breakpoint DNA.	2e-044
**************************************	**************************************	H.sapiens CpG island DNA genomic Msel fragment, clone	12C 077
2243	Z60212	195c8, forward read cpg195c8.ft1a	2e-045
****	M22485	Human ring chromosome 21 [r(21)] breakpoint DNA.	2e-045
·		Oryctolagus cuniculus peroxisomal Ca-dependent solute	<b>20-0-7</b>
2245	AF004161	carrier mRNA, complete cds	2e-045
	D38112	Human mitochondrial DNA, complete sequence	1e-045
		Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	10-0-3
2247	U72787	complete sequence [Homo sapiens]	6e-046
		Human DNA sequence from cosmid cN116A5, on	00-040
2248	Z69925	chromosome 22q12-qter contains exon trap	5e-046
·····	U12404	Human Csa-19 mRNA, complete cds.	5e-046
	Y07969	H.sapiens mRNA for APRIL protein	2e-046
\$1.0000 BUT COMMENCE CONTRACTOR AND AG	X90583	H.sapiens mRNA for rat translocon-associated protein delta	2e-046
	L35657	Homo sapiens (subclone H8 5_a10 from P1 35 H5 C8) DNA	
		Homo sapiens chaperonin containing T-complex subunit 6	20-0-0
	anni cana	(CCT6) mRNA > :: gb L27706 HUMTCP20 Human	
2253	NM 001762.1		2e-046
	AF008563		7e-047
***************************************	**************************************	Homo sapiens heterogeneous nuclear ribonucleoprotein	
		A2/B1 (HNRPA2B1) mRNA > ::	
2255	NM 002137.1		5e-047
***************************************		Homo sapiens cytochrome c oxidase subunit VIIb (COX7B),	30 01/
		nuclear gene encoding mitochondrial protein, mRNA > ::	
		emb Z14244 HSCOX7BM H.sapiens coxVIIb mRNA for	
2256	NM 001866.1	. ,	3e-047
-	**************************************	H.sapiens CpG island DNA genomic Mse1 fragment, clone	
2257	Z55938	1	2e-047
***************************************		Homo sapiens mitotic feedback control protein Madp2	
2258	U31278	· · · · · · · · · · · · · · · · · · ·	2e-047
***************************************	The state of the s	Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	
2259		·	2e-047
***************************************		Homo sapiens tumor protein, translationally-controlled 1	20-0-7
		(TPT1) mRNA > :: emb X16064 HSTUMP Human mRNA	
2260	NM 003295.1		7e-048
		Homo sapiens CDC7 (cell division cycle 7, S. cerevisiae,	, 0 0 10
		homolog)-like 1 (CDC7L1) mRNA, and translated products	
2261			3e-048
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	JU-040
2262			2e-048
			8e-049
2263	T119//		

SEQ ID	ACCESSION	DESCRIPTION	P VALUI
2265	M81600	Human NAD(P)H:quinone oxireductase gene, exon 6 >	2e-049
		Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	
2266	U72787	complete sequence [Homo sapiens]	8e-050
2267	M31004	Human nuclear phosphoprotein B23 mRNA, clone hpB2.	8e-050
2268	Z97207	Mus musculus mRNA for B-IND1 protein	8e-050
2269	AF007775	Rattus norvegicus aquaporin-pancreas and liver	3e-050
		Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	
2270	U72787	complete sequence [Homo sapiens]	9e-051
entration of the comment	\$1000000000000000000000000000000000000	Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	
2271	U72787	complete sequence [Homo sapiens]	9e-051
***************************************		Mouse mRNA for Cu-Zn superoxide dismutase (EC	<u> </u>
	98.	1.15.1.1) > :: gb M35725 MUSCZSOD Mouse Cu-Zn	And Andreas
2272	X06683	superoxide dismutase mRNA, complete cds.	9e-051
anachi diningana na katawa na manana na katawa na k	X93334	H.sapiens mitochondrial DNA, complete genome	3e-051
	U27197	Drosophila melanogaster pelota (pelo) mRNA, complete cds	
MARKET PROTOCOLOR DE LA COMPTION DE CONTRACTOR DE CONTRACT	D87953	Human mRNA for RTP, complete cds	1e-051
and the same of the same of the	L81869		4e-052
	D63876	Human mRNA for KIAA0154 gene, partial cds	4e-052
***********	D14659	Human mRNA for KIAA0103 gene, complete cds	3e-052
	X93334	H.sapiens mitochondrial DNA, complete genome	4e-053
entransamenta esta esta esta esta esta esta esta es	D38112		4e-053
	U97519	Homo sapiens podocalyxin-like protein mRNA, complete	4e-053
	L35657	Homo sapiens (subclone H8 5 a10 from P1 35 H5 C8) DNA	Same terretories and the second second
	D63876	Human mRNA for KIAA0154 gene, partial cds	4e-053
and consideration of the constraint of the const	D38112	Human mitochondrial DNA, complete sequence	1e-053
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	-
2285	Z57342	172a12, forward read cpg172a12.ft1a	1e-053
***************************************	**************************************	H.sapiens CpG island DNA genomic Mse1 fragment, clone	
2286	Z64479	127c4, reverse read cpg127c4.rt1a	1e-053
	M28209	Homo sapiens GTP-binding protein (RAB1) mRNA,	1e-053
**********	S57803	Ro60 protein gene [human, mRNA Partial, 176 nt, segment	4e-054
a produce accessor accessor for processor accessored	U01139	Mus musculus B6D2F1 clone 2C11B mRNA.	1e-054
		Homo sapiens cosmid clone U163C11 from Xp22.1-22.2,	
2290	U72787	complete sequence [Homo sapiens]	1e-054
	***************************************	Human tissue factor mRNA, complete cds, with an Alu	
2291	M16553	1	5e-055
	D10522	Homo sapiens mRNA for 80K-L protein, complete cds	5e-055
	Z71621	H.sapiens Wnt-13 mRNA	5e-055
***************************************	M81104	Human CD34 mRNA, complete cds.	4e-055
and the second	D29805	Human mRNA for beta-1,4-galactosyltransferase, complete	2e-055
***************************************		Human mRNA for liver alcohol dehydrogenase (EC 1.1.1.1)	
2296	X04299		2e-055
		Rattus norvegicus FGF receptor activating protein FRAG1	
2297	U57715	(FRAG1) mRNA, complete cds	2e-055
	iddele konsiliiriddaman.midirau.mar.gopeamaman.gopun magaruu.uu	Homo sapiens signal transducer and activator of	
land of the state		transcription 3 (acute-phase response factor) (STAT3)	
and the state of t		mRNA > :: gb L29277 HUMAPRF Homo sapiens DNA-	
2298	NM 003150.1	· · · · · · · · · · · · · · · · · · ·	2e-056
economical accommission and a continue of the	X56974	* <del>***********************************</del>	2e-050 2e-057
		Human nicotinic acetylcholine receptor alpha6 subunit	20-0 <i>3</i> 1
2300	U62435	1	2e-057
		H.sapiens CpG island DNA genomic Mse1 fragment, clone	20-031
	Z63454		6e-058

SEQ ID	ACCESSION	DESCRIPTION	P VALU
THE PARTY OF THE P	U34994		
2302	(U34994	Human DNA-dependent protein kinase catalytic subunit H.sapiens CpG island DNA genomic Mse1 fragment, clone	6e-058
2202	760422		2- 050
	Z60432	22h4, reverse read cpg22h4.rtla	2e-058
~~~	U83590	Rattus norvegicus PAR interacting protein mRNA, complete Rattus norvegicus PAR interacting protein mRNA, complete	
terroreiteraniano de contracto de la contracto	U83590 M28449	Mouse Hox-1.7 protein mRNA, 3' end.	and the comment of th
	M18981	Human prolactin receptor-associated protein	8e-059
2307	` [VII 0 9 0 I	Homo sapiens cytochrome c oxidase subunit VIIb (COX7B),	3e-059
		nuclear gene encoding mitochondrial protein, mRNA > ::	approximation and the second
		emb Z14244 HSCOX7BM H.sapiens coxVIIb mRNA for	
2308	NM_001866.1	cytochrome c oxidase subunit VIIb	2e-059
2300	,11111_001000.1	Homo sapiens tumor protein, translationally-controlled 1	26-039
		(TPT1) mRNA > :: emb X16064 HSTUMP Human mRNA	and the same of th
2300	NM 003295.1	for translationally controlled tumor protein	2e-059
***	D26067	Human mRNA for KIAA0033 gene, partial cds	9e-060
2310	D20001	Homo sapiens sorcin (SRI) mRNA > ::	76-000
2311	NM 003130.1	gb M32886 HUMSRICPA Human sorcin CP-22 mRNA,	9e-060
	X93334	H.sapiens mitochondrial DNA, complete genome	3e-060
nan Antonomoro menoro de la composición	U12404	Human Csa-19 mRNA, complete cds.	2e-060
	AF070661	Homo sapiens HSPC005 mRNA, complete cds	1e-060
	U77665	Human RNaseP protein p30 (RPP30) mRNA, complete cds	1e-060
	L03558	Homo sapiens cystatin B mRNA, complete cds.	9e-061
***************************************	D14048	Rat mRNA for SP120, complete cds	3e-061
***************************************	NM 002450.1	Homo sapiens metallothionein 1L (MT1L) mRNA >	3e-061
	annotatio var manaitra annotati considerativa annotati	Homo sapiens cytochrome c oxidase subunit VIIb (COX7B),	
		nuclear gene encoding mitochondrial protein, mRNA > ::	***************************************
		emb Z14244 HSCOX7BM H.sapiens coxVIIb mRNA for	
2319	NM 001866.1	cytochrome c oxidase subunit VIIb	4e-062
accompressed and the control of the	AC001443	Homo sapiens (subclone 2 f10 from BAC 2913	3e-062
		Homo sapiens heat shock 60kD protein 1 chaperonin	30 002
2321	NM 002156.1	(HSP60) mRNA, complete cds.	3e-062
	Z97207	Mus musculus mRNA for B-IND1 protein	3e-062
MONO/MANAGEM NOV. NOV.	J01415	Human mitochondrion, complete genome	1e-062
~~~~~	D83735	Homo sapiens mRNA for neutral calponin, complete cds	4e-063
	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Homo sapiens cytochrome c oxidase subunit VIIb (COX7B),	
		nuclear gene encoding mitochondrial protein, mRNA > ::	
		emb Z14244 HSCOX7BM H.sapiens coxVIIb mRNA for	
2325	NM 001866.1	,	4e-063
***************************************	X84694	H.sapiens mRNA for elongations factor Tu-mitochondrial	4e-063
	# /###################################	Homo sapiens cathepsin E (CTSE) mRNA > ::	
2327	NM_001910.1	gb J05036 HUMCTSE Human cathepsin E mRNA, complete	4e-063
and the second	AF007775	of particular and the contract of the contract	3e-063
2329	AF007775	Rattus norvegicus aquaporin-pancreas and liver	3e-063
2330	AF007862	Mus musculus mm-Mago mRNA, complete cds	3e-063
macominimum contrates and contrates	D38112	and the contract of the contra	1e-063
	***************************************	Homo sapiens succinate dehydrogenase complex, subunit D,	***************************************
one of		integral membrane protein (SDHD) mRNA > ::	
and a second		dbj AB006202 AB006202 Homo sapiens mRNA for	
2332	NM_003002.1	į	1e-063
	D38112		1e-063
	M12759	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1e-063
	U07802	Human Tis11d gene, complete cds.	1e-063

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SEQ ID	ACCESSION	DESCRIPTION	P VALU
2227	1120252	Rattus norvegicus initiation factor eIF-2B gamma subunit	064
	U38253	(eIF-2B gamma) mRNA, complete cds	6e-064
	M22146	Human scar protein mRNA, complete cds.	5e-064
2339	L20681	Rat proto-oncogene (Ets-1) mRNA, complete cds.	4e-064
		Human L1Heg repetitive element from the intergenic region	
2340	M14292	of the epsilon and G-gamma globin genes.	4e-064
		Homo sapiens tight junction protein 1 (zona occludens 1)	
_		(TJP1) mRNA > :: gb L14837 HUMZO1A Human tight	
2341	NM_003257.1	junction (zonula occludens) protein ZO-1 mRNA, complete	4e-064
		Homo sapiens succinate dehydrogenase complex, subunit D,	
		integral membrane protein (SDHD) mRNA > ::	
		dbj AB006202 AB006202 Homo sapiens mRNA for	
	NM_003002.1	cytochrome b small subunit of complex II, complete cds	2e-064
·····	X12883	Human mRNA for cytokeratin 18	2e-064
2344	D42044	Human mRNA for KIAA0090 gene, partial cds	1e-064
2345	Z97207	Mus musculus mRNA for B-IND1 protein	1e-064
		H.sapiens MRP RNA gene encoding the RNA component of	
2346	X51867	RNase MRP	7e-065
2347	X52104	Human mRNA for p68 protein	7e-065
2348	X74215	H.sapiens mRNA for Lon protease-like protein	6e-065
2349	U20796	Rattus norvegicus nuclear receptor Rev-ErbA-beta mRNA,	5e-065
2350	X79201	H.sapiens mRNA for SYT	5e-065
2351	U34584	Human Bcl-2 interacting killer (BIK) mRNA, complete cds	5e-065
		Human DNA sequence from cosmid U61F10, between	<u> </u>
2352	Z75894	markers DXS366 and DXS87 on chromosome X contains	4e-065
		Homo sapiens high-mobility group (nonhistone	<del></del>
authorized a		chromosomal) protein isoforms I and Y (HMGIY) mRNA >	Laws Johnson
2353	NM 002131.1	:: emb X14958 HSHMGY Human hmgI mRNA for high	2e-065
2354	AF007862	Mus musculus mm-Mago mRNA, complete cds	5e-068
2355	U74297	Oryctolagus cuniculus PiUS mRNA, complete cds	3e-069
orani terrarani terraran sendi,		Sus scrofa mRNA for soluble angiotesin-binding protein,	***************************************
2356	D11336	complete cds	2e-069
and the second s	AF007862	Mus musculus mm-Mago mRNA, complete cds	2e-069
	U74297	Oryctolagus cuniculus PiUS mRNA, complete cds	3e-071
~~~~~~~\$	AF061260	Mus musculus immunosuperfamily protein Bl2 mRNA,	5e-071
	U92949	Mus musculus kinesin motor protein KIFC2 mRNA,	2e-077
	AF007775	Rattus norvegicus aquaporin-pancreas and liver	6e-079
	AF007775	Rattus norvegicus aquaporin-pancreas and liver	8e-082
	U63840	Rattus norvegicus audiaporni-panereas and river	6e-085
	U57344		1e-085
annone programme province and a profession	Z11886		5e-087
en en et et den en e	AF007862	Mus musculus mm-Mago mRNA, complete cds	2e-089
2200	AF007862	Mus musculus mm-Mago mRNA, complete cds	1e-092
2367		M.musculus mRNA for map kinase interacting kinase, Mnk2	
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Y 1 1 1 0 9 7	intermediate increase for map kindse interacting kindse, Wilk2	
2368	~~~~ <del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	Gallys gallys exclin dependent bings (odles) cano complete	
2368	L77991	Gallus gallus cyclin-dependent kinase (cdk6) gene, complete	6e-098
2368 2369	L77991	Mus musculus fibroblast growth factor inducible gene 14	
2368 2369	~~~~ <del>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</del>	Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds	e-163
2368 2369 2370	L77991 U42386	Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds Mus musculus fibroblast growth factor inducible gene 14	e-163
2368 2369 2370	L77991	Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds	
2368 2369 2370 2371	L77991 U42386 U42386	Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds Mus musculus fibroblast growth factor inducible gene 14	e-163 e-160
2368 2369 2370 2371 2372	L77991 U42386	Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds Mus musculus fibroblast growth factor inducible gene 14 (FIN14) mRNA, complete cds	e-163 e-160 e-144

Table 2A	Nearest Neighbor (1	BlastN vs. GenBank)	
SEQ ID	ACCESSION	DESCRIPTION	P VALUI
2375	AJ000696	Rattus norvegicus mRNA for a novel kinesin-related protein,	e-101
2376	Z97207	Mus musculus mRNA for B-IND1 protein	e-102
2377	AJ000696	Rattus norvegicus mRNA for a novel kinesin-related protein,	e-122
2378	AB000172	Porcine mRNA for endopeptidase 24.16, complete cds	e-118
2379	AB000171	Porcine mRNA for endopeptidase 24.16, complete cds	e-131
		Rattus norvegicus initiation factor eIF-2B gamma subunit	and a second
2380	U38253	(eIF-2B gamma) mRNA, complete cds	e-129
2381	X54352	M.domesticus MD6 mRNA	e-142
2382	X14678	Mouse TPA-induced TIS11 mRNA	e-121
2383	X82632	M.fascicularis mRNA for NAD+-isocitrate dehydrogenase	e-142
2384	U42385	Mus musculus fibroblast growth factor inducible gene 16 (FIN16) mRNA, complete cds	e-123
2385	U17901	Rattus norvegicus phospholipase A-2-activating protein (plap) mRNA, complete cds.	e-116
2386	U08215	Mus musculus Hsp70-related NST-1 (hsr.1) mRNA,	e-119
2387	X82632	M.fascicularis mRNA for NAD+-isocitrate dehydrogenase	e-148
2388	U63840	Rattus norvegicus nucleoporin p54 mRNA, complete cds	e-131
2389	U42385	Mus musculus fibroblast growth factor inducible gene 16 (FIN16) mRNA, complete cds	e-145
	U42385	Mus musculus fibroblast growth factor inducible gene 16 (FIN16) mRNA, complete cds	e-149
2391	U81045		e-114
2392	L77991	Gallus gallus cyclin-dependent kinase (cdk6) gene, complete	e-103
2393	U81045	rake, was war was a war and the same and the	e-115
2394	D30666	Rat mRNA for brain acyl-CoA synthetase II, complete cds Rattus norvegicus phospholipase A-2-activating protein	e-139
2395	U17901	(plap) mRNA, complete cds.	e-135
	D30666	Rat mRNA for brain acyl-CoA synthetase II, complete cds	e-126

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1	734646				l	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) (Colon Tumor Tissue	-	<b>-</b>		
i		vs. Colon Metastasis)	14	0	14.22	l
		15,16 (Normal Colon vs. Colon Tumor Tissue) (Normal Colon vs.				
		Colon Tumor Tissue)	0	14		13.25
2	400221					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	2	6.87	<del></del>
3	205329	<del> </del>				
		<del> </del>			<del> </del>	<del> </del>
		15.16 (Normal Colon vs. Colon Turnor Tissus)	178	7	26.88	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1/8		20.88	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis) (Normal Colon Tissue vs. Colon Metastasis)	178	1	191.06	ļ
		18,19 (Normal Colon Tissue vs. Colon Tumor) (Normal Colon Tissue	1/8	'-	191.00	<b> </b> -
		vs. Colon Tumor)	21	0	24	Ì
		18,20 (Normal Colon Tissue vs. Colon Metastasis) (Normal Colon	-21	-	24	<del> </del> -
		Tissue vs. Colon Metastasis)	21	0	17.95	į.
4	446680	1133uc vs. Colon Metastasis)	21	-	17.93	<del> </del>
	++0080	<del> </del>	<del> </del>	<del> </del>	<del></del>	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	29	84		2.7
-		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue) (Normal Lung				
		Tissue vs. Lung Tumor Tissue)	40	94	ļ	2.33
5	1261					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
6	400258					
			<del>                                     </del>			
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
7	450559	(Colon Fund Colon Michaelle)	۰	Ť	0.05	ļ——
	450555					
		1(17/0-1 - T T 0 1 1 1 1 1 1 1 1	<u> </u>			5.40
	150050	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	11		5.42
8	450959					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	9		8.51
9	451794					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	0	15.85	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	1	16.1	
10	415058	<del> </del>				
		<del> </del>				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	-		5.01
<del>-,, -  </del>	21500	10,17 (Colon Tunior Lissue vs. Colon Metastasis)	0	6		5.91
11	31506	<b> </b>				
l		15,16 (Normal Colon vs. Colon Tumor Tissue)	20	77		3.64
		Low Met)	5	0	6.99	
1		15,17 (Normal Colon Tissue vs. Colon Metastasis)	20	58		2.7
12	417155					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
13	448925	2012 (Committee Colon Fullor House)		<del></del> -	0.57	
	T7074J					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	5	15		2.95
14	11329			1	1	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		10.20 (G.1., T, T, G.1.)	<u> </u>			ļ
		19,20 (Colon Tumor Tissue vs. Colon Metastasis) (Colon Tumor Tissue vs. Colon Metastasis)	30	5	4.49	1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	112	38	3.12	<del> </del>
15	650422				<b> </b>	<del> </del>
						<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	18	0	19.32	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	18	6	3.17	
16	6863					
			ļ	ļ	ļ	
		01,02 (Colon, High Met vs. Colon, Low Met)	1	8	<b></b>	8.67
17	449690			<b>}</b>	}	
		1612/01 7	<del> </del> _			
18	724616	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	17		5.58
18	724616				}	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8	<del> </del>	7.57
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	7.57
19	549722	1317 (COOR Tariot 115550 15. COOR Metastasis)	Ť	ĻŤ	0.12	<del> </del>
		<u> </u>				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
20	549722					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
21	448110					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	25		11.65
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	25		24.62
22	515631					
		151(0)				
23	11001	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
	11881					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
24	650856	55,64 (Blodd, 11gh Met 15. Blods, 16h Met)		<u> </u>	5.05	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
25	449701					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	17	1	17.26	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	17		16.08
26	651073					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
	10240	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
27	10340					
		03,04 (Breast, High Met vs. Breast, Non-Met)		0	505	
28	648310	out, of Chicast, right vict vs. Dicast, Non-Met)	6	<del></del>	5.85	
	040310					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
29	730336	· · · · · · · · · · · · · · · · · · ·				

Table 5

Table 5		T. Sharama Dalan A. D		T D	1 A /D	I D/A
SEQ	CLST	Library Pair A,B	A	B	A/B	B/A
		15.14 (Namuel Colon via Colon Turner Tiesus)		24	<del> </del>	22.71
		15,16 (Normal Colon vs Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	24	0	24.37	22.71
30	3060	10,17 (Colon Tunior Tissue vs. Colon Wetastasis)		<del>                                     </del>	24.31	<del> </del>
	3000			<del> </del>	<del> </del>	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	17		4.19
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	4	3.7	7.12
		03,04 (Breast, High Met vs. Breast, Non-Met)	17	11	·	11.27
31	453016	(		<del>                                     </del>		11.27
					·	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	1
33	185461			<u> </u>		
				<del>                                     </del>		
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	17		17.42
34	452530		<del></del>			
	<del></del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
35	448925			-		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	5	15		2.95
36	1013					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		01,02 (Colon, High Met vs. Colon, Low Met)	40	84		2.28
37	6545					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
38	449891					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8 46	
39	4045					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	11		5.96
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	
40	404475					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	2	5.59	
-47	650307	15,17 (Normal Colon Tissue vs. Colon Metastasis)	19	2	10.2	
41	650297					
		15 17 (Normal Colon Tions vs. Colon Material)		<u> </u>	6.44	
42	650493	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
72	030493					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
43	644884	15,17 (110111111 COIOII 11050C vs. COIOII INCIASIASIS)	<del></del>		7.31	
	01-100-1					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
44	452212	Colon Memorial		<del></del>	0.05	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
	402727	<del></del>				

Table 5

-CE	CY COR	T		1 -		T = 1.1
SEQ	CLST	Library Pair A,B	+A	В	A/B	B/A
		151701 101 7	+	<del> </del>	<b> </b>	
46	645104	15,17 (Normal Colon Tissue vs. Colon Metastasis)	5	17		3.17
46	645194		+	}		<del>}</del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<del> </del>
47	447501	13,17 (Northal Colon Tissue vs. Colon Metastasis)	+	-	7.31	<del> </del>
<del>- 7</del> /	447301			<del> </del>		├
<del></del>	<del></del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	3	5.37	<del> </del>
48	556326	1537 (Frontial Colon Nasac vs. Colon Newstasis)	+		3.37	╁
<del>;°</del>		<del> </del>	+			<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	8		7.88
49	447035		† <u> </u>	<u> </u>		1.00
			+			<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	<del> </del>
50	2551		+- <u>-</u> -	<del> </del>		<del> </del>
			<del> </del>	-		<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<del> </del>
51	736154		1			<del> </del> -
						<b></b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
52	452028		1			
			1			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
53	447441		1			
			1			i
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	34	129		3.53
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	34	129		3.74
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		10.7
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	155	32	4.89	
54	11028					
}		01,02 (Colon, High Met vs. Colon, Low Met) (Colon, High Met vs.				
	640074	Colon, Low Met)	0	6		6.5
55	640974	<u> </u>	<del></del>			<b> </b> -
		15.17 (Named Colon Tissue vs. Colon Metastasis)	+		0.66	<b></b> -
56	555103	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	<u> </u>
	333103		+			<del></del>
}		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	6		5.94
57	446789	25324 (Normal Collig 1188de VS. Lulig Tullior 1188de)	+			3.94
		<del></del>	+			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	5	3.38	
58	644884		+-~			
			+			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
59	9029		1			
			1			
		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	
60	419255					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	0	11.63	

Table 5

SEQ	CLST	Library Pair A,B	A	B	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	1	11.81	T
61	4309					
						<b>†</b>
		01,02 (Colon, High Met vs. Colon, Low Met)	4	13		3.52
62	554069		1	1	<b></b>	1
			1	<del>                                     </del>		<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	6		5.91
63	4330	10,17 (COION TURNOT TISSUE VS. COION TVCULSUUSIS)	+			3.71
-03	4330		+			<del> </del>
		03,04 (Breast, High Met vs. Breast, Non-Met) (Breast, High Met vs.	+			
- 1		Breast, Non-Met)	13	3	4.23	]
		01,02 (Colon, High Met vs. Colon, Low Met)	1 1	10	7.23	10.84
64	644903	77,02 (Colon, Tigh Net vs. Colon, Low Met)	<del>                                     </del>	10		10.04
	044903	<u> </u>	<del> </del>			<del> </del>
		15.17.01	+			
	710005	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	ļ
65	549395					<b></b> _
<del></del> l				ļ		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	13	L	6.06
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	13		12.8
66	4974					
1						
		01,02 (Colon, High Met vs. Colon, Low Met)	1	8		8.67
67	447466		T			
			T			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	155	5	32.77	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	16	1	18.28	<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	155	2	83.19	<del>                                     </del>
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	16	0	13.68	<del>                                     </del>
68	645073	(1 to final colon fissac to. colon fiscassasis)	+	Ť	15.00	<del> </del>
	043073		+			<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<del></del>
		<del>                                     </del>	<del></del>			<del> </del>
	447070	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
69	447978		<del> </del>			ļ
			1			ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
70	607430		1	igsquare		
					·	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	L
71	556198		1			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
72	450323		$\perp$			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
73	21205					
	<del></del>					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	9		9.22
74	561109	<del> </del>				<del></del>
	<del></del>	<del> </del>	+			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
75	446673	1970 (Assimal Colon vs. Colon Taillot 11880C)	+			1.57
	440073	<del> </del>	<del> </del>			<b> </b>
<del></del>						
15		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	

Table 5 SEQ CLST Library Pair A,B A В A/B B/A 15,16 (Normal Colon vs. Colon Tumor Tissue) 24 4 6.34 15,17 (Normal Colon Tissue vs. Colon Metastasis) 24 3.22 8 77 449142 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 9 1 9.14 78 5830 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 7 0 7.11 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 6.62 554109 79 15,16 (Normal Colon vs. Colon Tumor Tissue) 15 2 7.93 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 2 11 5.42 80 595506 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 6 5.68 453981 81 23,24 (Normal Lung Tissue vs. Lung Tumor Tissue) 43 258 5.94 19,20 (Colon Tumor Tissue vs. Colon Metastasis) 2 110 73.53 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 590 99.86 6 15,17 (Normal Colon Tissue vs. Colon Metastasis) 68 12.16 6 15,16 (Normal Colon vs. Colon Tumor Tissue) 68 590 8.21 18,20 (Normal Colon Tissue vs. Colon Metastasis) 110 0 128 69 82 642461 15,17 (Normal Colon Tissue vs. Colon Metastasis) 14 0 15.03 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 10 0 10.16 83 556198 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 6 5.68 84 2082 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 6.62 03,04 (Breast, High Met vs. Breast, Non-Met) 15 55 3.76 08,09 (Lung, High Met vs. Lung, Low Met) 7 38 3.88 85 549435 15,16 (Normal Colon vs. Colon Tumor Tissue) 14 2 7.4 2286 86 15,16 (Normal Colon vs. Colon Tumor Tissue) 5 28 5.3 19,20 (Colon Tumor Tissue vs. Colon Metastasis) 13 2 4.86 15,17 (Normal Colon Tissue vs. Colon Metastasis) 5 19 3.54 18,19 (Normal Colon Tissue vs. Colon Tumor) 3 13 3.79 2737 87 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 3 14 4.6 15,17 (Normal Colon Tissue vs. Colon Metastasis) 4 14 3.26 88 728115 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 7 6.62

7

0

711

16,17 (Colon Tumor Tissue vs. Colon Metastasis)

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
89	650856					
	ļ					1
	ļ	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
90	650476					
	<u> </u>					
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
91	535208					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
92	733849					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
93	447978					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
94	729483					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
05	12010	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
95	12018			<u> </u>		
		00.04.79		<u> </u>		
-06	42.45	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
96	4747					
		01.00 (0.1				
97	4747	01,02 (Colon, High Met vs. Colon, Low Met)	2	10		5.42
91	4747					
		01.02 (C.1., W.1.)		<u> </u>		
98	185577	01,02 (Colon, High Met vs. Colon, Low Met)	2	10		5.42
76	105577					
		03 04 (Propost High Motors Propost N. 14.0)				
99	4126	03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2
-	.120					
<del></del>		03,04 (Breast, High Met vs. Breast, Non-Met)		-		
100	11456	object (Bleast, High Wet vs. Bleast, Non-Wet)	13	1	12.68	
		01,02 (Colon, High Met vs. Colon, Low Met)				
101	729851	( Colon, Mg. Met 13. Colon, Low Met)	0	6		6.5
T T		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	711	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	7.11	( (2
102	449849	( 1888)	<del>-   °  </del>			6.62
				-+		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.01
103	2490		<del></del>	- +		5.91
			<del>-  -  </del>			
		08,09 (Lung, High Met vs. Lung, Low Met)	7	- +	9.78	
		01,02 (Colon, High Met vs. Colon, Low Met)	21	6	3.23	
		03,04 (Breast, High Met vs. Breast, Non-Met)	14	3	4.55	
104	549041				7.33	
				-+		
	1	6,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91

SEQ	CLST	Library Pair A,B	T A	В	A/B	<b>B</b> /A
105	11881			†		1
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
106	724296			ļ		
	<u> </u>	15,16 (Normal Colon vs. Colon Tumor Tissue)		1.25		<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	128	128	120.00	121
107	726173	(Colon Turner Fissac Vs. Colon Micrastasis)	128	0	129.99	
						<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
108	2423					
		03.04 (D				
109	556250	03,04 (Breast, High Met vs. Breast, Non-Met)	14	0	13.66	
107	330230				<del></del>	
	<u>.                                    </u>	15,16 (Normal Colon vs. Colon Tumor Tissue)	1	28		26.4
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	28	4	7.11	20.4
110	643594				7.11	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
111		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
111	11881					
		03,04 (Breast, High Met vs. Breast, Non-Met)			5.0.5	
112	7436	55,64 (Bleast, High Met vs. Bleast, Non-Wet)	6	0	5.85	
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
113	2110					
114	10240	03,04 (Breast, High Met vs. Breast, Non-Met)	31	5	6.05	
114	10340					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.05	
115	643594	oos, (Sound, Ingh Met vs. Bleast, Ivon-Wet)	- 6	-	5.85	
				-+		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
116	447035					
		16.17 (Color Turner Times Color				
117	402707	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	
-11	702,07					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.38
118	645799	, (Control of Control	1	<del>-                                    </del>		0.30
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
119	171511					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
120		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
120	451607			_		
		6,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B
121	3138					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	10		10
122	2988			<u> </u>		<u> </u>
				<u> </u>		<u> </u>
		01,02 (Colon, High Met vs. Colon, Low Met)	2	11		5.
123	447326			-		<u> </u>
		15 17 Olympia Calar Time or Calar Vista (a)			C 44	╂
124	561734	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	$\vdash$
124	301734			-		╁
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	<del> </del>	5.
125	454999	10,10 (Activation Colon Value Planter Planter)	Ť	Ť	<u> </u>	+ -
				-		<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	28	15	2	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	28	11	2.69	
126	185652					
						L
		03,04 (Breast, High Met vs Breast, Non-Met)	0	7		7.
127	6725					
				<u> </u>		_
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	<u> </u>
128	726644					_
		151(2) 151 21 7		10		<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	18	10.70	17
129	11012	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	18	0	18.28	
129	11012					-
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6
130	726377					۳
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.
131	735326					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.
122	650015	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	_
132	650845					<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.00	$\vdash$
133	9048	10,17 (Coloit Tuthor Tissue vs. Coloit Metastasis)	0	U	6.09	
	20-10					$\vdash$
<del></del>		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	$\vdash$
134	732254	. ,		-	2.70	$\vdash$
						$\vdash$
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.0
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	8		7.9
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
135	452052					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
136	554079					
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	2	10		4.9

Table 5 Page 9 of 155

SEQ	CLST	Library Pair A,B	A	В	A/B	
137	9049					Γ
		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	L
138	1307			<u> </u>		L
		100 04 (P		- 00		╀
120	120720	03,04 (Breast, High Met vs. Breast, Non-Met)	14	89		╀
139	139730					╁
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		H
140	7750					T
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	14		]
141	8050			ļ		L
<u> </u>				_		L
142	725222	01,02 (Colon, High Met vs. Colon, Low Met)	0	7		╀
142	123222			<b></b>		╁
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		Η,
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
143	3275					
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	0	7.81	L
144	7424	- Alaba				ـ
		02 04 (Decet High Maters Decet New Mat)		_	5.05	╀
145	8953	03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	$\vdash$
143	6733					H
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	13		1
146	8966					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		Ľ
147	530883			$\vdash$		├
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		۱,
148	6725	13,10 (Normal Colon vs. Colon Turnor Tissac)				<del>  `</del>
						-
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	Ī
149	4439					$\Box$
		00.00 (I H' 1 M )			6.00	$\vdash$
		08,09 (Lung, High Met vs. Lung, Low Met) 01,02 (Colon, High Met vs. Colon, Low Met)	10	2	6.99	$\vdash$
150	648472	or,02 (Colon, Tright Meet vs. Colon, Low Met)	13		0.92	$\vdash$
	0.0472			H		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	H
151	735346					Г
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		
153		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
152	732121			$\vdash \vdash \vdash$		<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	-
ı						

SEQ	CLST	Library Pair A,B	A	B	A/B	]
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	Т
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
154	533588					T
						1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	$\top$
155	649667					$\vdash$
100	0.12001					$\vdash$
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	╁
156	394436	13,17 (Normal Colon Hissac Vs. Colon Medicals)	<u>~</u>	-	0.44	╁
150	394430			-		╁
		15 17 (Namuel Colon Tienne un Colon Metestoria)	7	0	7.51	╁╌
1.57	(40254	15,17 (Normal Colon Tissue vs. Colon Metastasis)	<del>-                                    </del>	0	7.51	-
157	649354			-		
		15.15.01		-		-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	┼
158	2022					-
						↓_
		03,04 (Breast, High Met vs. Breast, Non-Met)	5	15		3
		21,22 (Normal Prostate vs. Prostate Cancer)	0	6		<u> </u>
159	561359					$oldsymbol{oldsymbol{\perp}}$
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	3	4.29	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	2	6.34	
160	7607	"				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	3	4.65	
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	4	3.44	
161	7750					
						T
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	14		14
162	410554					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs Colon Tumor Tissue)	0	6		5
163	2315	·	1			1
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	118		11
		01,02 (Colon, High Met vs. Colon, Low Met)	25	4	5.76	П
164	561734					T
				$\vdash$		T
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5
165	4420	, , ,				۲
	•			$\vdash$		$\vdash$
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	$\vdash$
		01,02 (Colon, High Met vs. Colon, Low Met)	1	10	5.05	10
166	559663	5 1,02 (Colon, riigh frict vs. Colon, Low Mct)	1	''		<del>  '</del>
100	227003			<del>                                     </del>		├
		15 16 (Namuel Cales et Cales Transcort'			2.06	$\vdash$
1/7	7000	15,16 (Normal Colon vs. Colon Tumor Tissue)	15	4	3.96	-
167	7082					-
				$\vdash$	0 = -	
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0 [	9.76	1

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	118		17.28
		01,02 (Colon, High Met vs. Colon, Low Met)	25	4	5.76	17.20
169	650472					
170	(492	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
170	6482					<del> </del>
	<del></del>	01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
171	4584					
		01,02 (Colon, High Met vs. Colon, Low Met)	1	11		11.93
172	453846	or,oz (Colon, riigii Met 13. Colon, Low Mec)		111		11.5
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	11		10.25
173	650820					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
174	642906					
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	7	0	7.11	
175	448805	10517 (Colon Tullior Fissac vs. Colon Metastasis)		Ť	7.11	
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	6	20		3.28
176	649667					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
177	735786					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	,,,,,	6.62
178	121457					
		00.00 (1 Hish Mans Law Law Ma)	01	359		2.82
		08,09 (Lung, High Met vs. Lung, Low Met) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	91	11		5 12
179	372960	Total (Average of the Color Matter and the Color Ma				
				10		2.79
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	6	18 33		5.2
180	120049	12310 (100mm) Colon Vs. Colon Tumor Tusse)		55	*****	
101	649006	15,17 (Normal Colon Tissue vs Colon Metastasis)	10	2	5.37	
181	648996			$\vdash$		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
182	3765					
		01,02 (Colon, High Met vs. Colon, Low Met)	19	6	2.92	
183	462642					
	9	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	Ð	6.34	ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<del>                                     </del>

SEQ	CLST	Library Pair A,B	A	В	A/B	B
184	727181					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
185	649259					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	$\vdash$
186	649717	13,17 (Normal Colon Tissue vs. Colon Metastasts)		ľ	7.51	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	_
187	736860			-		-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.
188	729175					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	_
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.09	5.
189	642906	15,10 (NOTHER COION VS. COION TUMOr HISSUE)	- <sup>0</sup>	"		-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
190	4420					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	-
		01,02 (Colon, High Met vs. Colon, Low Met)	1	10		10
191	2420					
		Ol Ol (Calar Wat Many Calar Law Ma)	20	11	2.35	<u> </u>
192	648109	01,02 (Colon, High Met vs. Colon, Low Met)	28	11	2.33	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1_	8.46	_
193	2334					├
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.
194	639705					
		15 16 Oleveni Cales and Cales Town Time	10		10.57	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.57	<del> </del>
195	551907	15,17 (NOTHIAL COLOR TISSUE VS. COLOR METASTASIS)		۲	10.73	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
196	561382					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	2	6.34	-
197	595506					
198	499424	15,16 (Normal Colon vs. Colon Tumor Tissue)	- 0	6		5.
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.
199	735477	l		[ ]		<u></u>

SEQ	CLST	Library Pair A,B	A	В	A/B	B/
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.
200	734370					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	11		10
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	10
201	779	10,17 (Color Turior 113de vs. Color Metasasis)		Ť,	11.17	$\vdash$
		01,02 (Colon, High Met vs. Colon, Low Met)	27	54		2.
202	649143	03,04 (Breast, High Met vs. Breast, Non-Met)	60	22	2.66	├-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	Ĺ
203	489					ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	58	<del></del>	4
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	58		4.
204	2994					
		01,02 (Colon, High Met vs. Colon, Low Met)	3	14		5.
205	2994	or,oz (Colon, Tiigh Met vs. Colon, Low Met)		14		3.
		01,02 (Colon, High Met vs. Colon, Low Met)	3	14		5.
206	11147					
-		01,02 (Colon, High Met vs. Colon, Low Met)	0	6	·	6
207	549395					
				12		
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	13		12
208	559806	to, i (Colon Tumor rissue vs. Colon Metastasis)		13		12
209	452238	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.
209	432238					<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.0
210	225914					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
211	463480	10,17 (COOR TERMS 11350C V3. COOR MCCastasis)	† <u>*</u>	Ť		
212	104727	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	0	6		5.9
212	184725					-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
213	557401			<del>                                     </del>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
214	455155					
215	551117	15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	

SEO	CLST	Library Pair A,B	A	В	A/B	B/A
3 <u>2 Q</u>						
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
217	729295					
	7.2.2.2.2					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.6
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
218	450429					
210	130 125					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	
-		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	1	13.95	
219	450148					
217	150110					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
220	380412	10,17 (COION TONIO 15000 157 COION				
220	360412					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
221	446614	15,10 (Ivormat Colon vs. Colon value Tiesas)				
221	440014					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
222	555911	10,17 (Colon Tullior Fissac vs. Colon Metastass)				<u> </u>
222	333911			$\vdash \vdash$		T
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	21		9.3
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	21		20.
222	450000	10,17 (Colon Turnor Tissac vs. Colon Metastasis)				$\vdash$
223	450828		-			<b> </b>
	-	15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.3
224	28	15,17 (Normal Colon Tissue vs. Colon Metastasis)				
224	2.0					$\vdash$
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	2	11		6.4
		18,19 (Normal Colon Tissue vs. Colon Tumor)	2	43		18.
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	53	207		3.0
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	697	1789		2.
	<u> </u>	19,20 (Colon Tumor Tissue vs. Colon Metastasis)	43	11	2.92	t
225	446450	19,20 (Cololi Tullioi Tissue vs. Cololi Metastasis)				1
225	440430			$\vdash$		<b>†</b>
	ļ	15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3.88	
226	452026	15,10 (Normal Colon vs. Colon Tumol Tissue)				
226	432020					
	<del>                                     </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	35	14	2.64	
227	643594	10,10 (Admin Colon 15. Colon Lamor Alouae)				
221	043374			1	-	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	<b>T</b>
	+	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	1
220	1905	13,17 (1401Hiai Cololi Tissuc vs. Cololi McLastasis)		Ť		1
228	1903			$\vdash$		
		01,02 (Colon, High Met vs. Colon, Low Met)	7	21		3.
	-	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.
220	651072	13,17 (NOITHAL COIGH TISSUE VS. COIGH MELASIASIS)	<del>-                                     </del>	t		† <u> </u>
229	651073			+		1
		15.17 (Normal Color Tionyo v. Color Metastage)	7	0	7.51	+
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.4	+
	1	15,16 (Normal Colon vs. Colon Tumor Tissue)	<del></del>	+-	- ' · <del>-</del>	+
230	553705			+		

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	0	12.68	
231	521840					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
232	648689					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
233	447858					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	
234	556198					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
235	394436					
	•					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
236	639651					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
237	499424					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
238	468109					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
239	185701					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
240	451811					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
241	730670		<u> </u>			
					,	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
242	172013					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
243	449142					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	- 1	9.14	
244	446964					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32
245	414739					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	14		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	14		6.52
246	641124					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
247	555702					
į.		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
248	549435					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	2	7.4	
249	643954	15,16 (Normal Colon vs. Colon Turnor Tissue)	14	2.	7.4	<u> </u>
247	043234					<b></b>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
250	5984					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
251	560526			-		
	-	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.8
252	411113	10,17 (Coloii Tunioi Tissue vs. Coloii Metastasis)		Ů		7.0
	111113					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
253	7607					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	4	3.44	-
		01,02 (Colon, High Met vs. Colon, Low Met)	13	3	4.65	7.5
	559409	15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	3	4.03	
	339409					-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
255	650053					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	1	10.57	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	ļ
256	448511			<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	26		8.0
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	26		6.4
257	642142					
			-		•	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	11		5.2
258	470462					
		15 17 Ol and Other Times of Color Materials	7	0	7.51	<u> </u>
259	431601	15,17 (Normal Colon Tissue vs. Colon Metastasis)		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	7.31	
260	431001			<b></b>		
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	9	0	7.69	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	28	9	3.34	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	28	6	4.93	
		08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	
	421431					ļ
	-	15 16 Oleman Colon via Color Transa Tirana	-	-	Q 16	ļ
261	284586	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
201	204380		<del> </del>			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
262	556198					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.6
263	431601					

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SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
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_		18,20 (Normal Colon Tissue vs. Colon Metastasis)	9	0	7.69	
		08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	28	6	4.93	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	28	9	3.34	
264	449891					
					0.16	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
265	556561					
		15.17 Ol Colon Tiggue ve Colon Metastasis)	0	10		9.32
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 16,17 (Colon Turnor Tissue vs. Colon Metastasis)	1	10		9.85
266	554188	10,17 (Colon Tunior Tissue vs. Colon Victasiasis)				
200	334100					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
267	3247					
		01,02 (Colon, High Met vs. Colon, Low Met)	1	20		21.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	26		3.66
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	26		6.06
268	546705					
						5.91
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		3.91
269	560984					
		16.17 (Color Times Tiesus vs. Color Motostosis)	0	6		5.91
270	455020	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	<del></del>			
270	455820					-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
271	643129	10,17 (2010) 11111				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
272	454653					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	17		5.28
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	5	17		3.35
273	456549					<del>                                     </del>
		Caller Material	1	8		7.88
	151005	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	0		7.00
274	454806					<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
275	724296	13,10 (Normal Colon vs. Colon Tumor 113340)				
213	724270					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	128		121.
	<b>+</b>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	128	0	129.99	
276	559280					
	1					<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
277	171511			<b> </b>		<b>_</b>
				<u> </u>		<del>                                     </del>
	I	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	<b>L</b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62

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Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	ļ
279	734370					
		15.16 (Names) Colon vs. Colon Tymos Trayya)	0	11		10.41
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	0	11.17	10.41
280	639459	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
280	033433					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
281	641679					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
282	644611					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
283	550038					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
284	452567					
		0.1.10				16
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	21	14	7.4	4.6
205	411113	15,16 (Normal Colon vs. Colon Tumor Tissue)	Z1	3	7.4	
285	411113					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
286	650749	1011 (Colon Tunior Tissue 151 Colon Memorials)				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
287	558899					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
288	452986					
						ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	<b> </b>
289	393197					
		16 17 (Color Transport Color March 19	<del></del>		77 1 1	
200	100404	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
290	499424			$\vdash$		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8	· <del></del>	7.57
291	21669	15,10 (Normal Colon vs. Colon Tumor Tissue)	- <u>-</u>			1.57
٠,٠١	21007					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
292	640590					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
		13,10 (Normal Colon vs. Colon Tunior Tissue)			7.31	<u></u>

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
294	448770					
				<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
295	559280					
	<u> </u>			<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
296	648934					
		151601 101 01 0		<u> </u>		
	ļ	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	ļ
207	152695	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	ļ
297	452685			<b> </b>		<u> </u>
		15.16 (Normal Colon va Colon Turnor Tiggue)	15		2.17	1
298	456549	15,16 (Normal Colon vs. Colon Tumor Tissue)	15	5	3.17	
290	430349			ļ	-	<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		8		7.00
299	446614	10,17 (COIOII TUINOI TISSUE VS. COIOII IVIETASTASIS)	1		ļ	7.88
200	140014				<u> </u>	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
300	559280	10,11 (Cotoli Tanot Tissae Vs. Cotoli Metasasis)		<del>                                     </del>	7.11	<del>                                     </del>
			· · · · · · · · · · · · · · · · · · ·			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
301	446673			<u> </u>		5.02
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	
302	562550					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
303	467288					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
304	463824					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
20.5	202107	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
305	393197					
		16.17 (Color Tymor Tierro Golor Mary				
306	407077	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
300	407077					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	1	8		7.45
307	499424	1317 (Colon Lunor Fissue Vs. Colon Michaelasis)	1	- 0		7.88
	122.12.1					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
308	554500	2770 (137116) COION 10. COION 10110/1 11350C)	<del></del>	0		1.31
			<del>-   -  </del>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
309	730143	7	<u>-</u> -	<del>-</del>	U.J.T	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		,				

Table 5

SEQ	CLST	Library Pair A,B		В	A /D	1D / A
310	595506	Library Fair A,B	A	В	A/B	B/A
310	3,3300					<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	<b></b>	5.68
311	2334	Table ( Table )		†	<del></del>	3.00
		***************************************		$\vdash$		†
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
312	647444					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
313	380291					
<u> </u>						
<u></u>		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
314	644849			L		
ļ						
<u></u>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	<u> </u>
315	449457					
		15 17 0/2 1 Color T'				
316	446673	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7	ļ	6.52
310	4400/3		<del> </del>			
	ļ	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	0 12	
317	549069	10,17 (Colon Turnor Tissue vs. Colon Metastasis)	-   °	1	8.12	ļ
	347007					
l		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	56		4.82
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	36		3.05
318	728884					3.00
					-	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
319	415058					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
320	553955					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
321	455820	d with				
<u> </u>		16.17.60.1 T. T. O.1. M.				
322	549617	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
322	349017					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5 60
323	449831	125,10 (Arothur Colon 15. Colon Tunior Tissue)	0	0		5.68
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	0	10		9.85
324	451580		<del>-   *  </del>			7.55
						-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	.=	5.91
325	558899					
	-					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
326	562292					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
327	5830					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
				ļ		
·		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	7.11	6.62
328	8953	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
	0,55					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	13		13.32
329	8012					
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
330	185718					
		Of OA (Decree With Mark Decree With Mark				
331	729851	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
331	729031					<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	7.1.2	6.62
332	185597					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2
333	9887					
224	72.502.5	03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
334	725825					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	1.51
335	6545	1 South Lames Albade 151 Colon Metablasis		Ů	0.12	
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
336	21205					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	9		9.22
337	8867				:	
		03,04 (Breast, High Met vs. Breast, Non-Met)			£ 9.5	
338	729295	05,04 (Bleast, High Met vs. Bleast, Non-Met)	6	0	5.85	
330	12,2,3					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	1	6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
339	730430					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
240	7070	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
340	7072					<del></del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	
341	730533	25,5 . (Stoudy Light Mot 15. Diedst, Holl-Mct)	10	1	7.10	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
342	9121					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59
343	11131					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
					ļ	
344	640116	01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
311	040110			1-		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	4	5.02	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	19	0	20.39	
345	730282					
				ļ		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	ļ
346	550571	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
340	330371			<del> </del>		<u> </u>
	<b> </b>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
347	1183			Ť		3.51
				<b>1</b>		
		01,02 (Colon, High Met vs. Colon, Low Met)	4	71		19.24
		03,04 (Breast, High Met vs. Breast, Non-Met)	32	15	2.08	
348	449437					
		15 16 Olympic 1 G 1		<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	14	3	4.93	2.04
349	8966	10,17 (Colon Tunior Tissue vs. Colon Metastasis)	3	12		3.94
	0,00			-	<u> </u>	
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
350	6134					
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	22	5	3.76	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	142	40	3.81	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	142	42	3.57	ļ
351	95700	19,20 (Colon Tumor Tissue vs. Colon Metastasis)	22	5	3.29	
-	70,00		· · · · · · · · · · · · · · · · · · ·	-	*, ,	
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	21		21.52
352	7066				·-··	
		01,02 (Colon, High Met vs. Colon, Low Met)	0	9		9.76
353	648310					
-	an	15,17 (Normal Colon Tissue vs. Colon Metastasis)				
$\dashv$		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.44	
354	730059	2000 (Totalia Colon Vii Colon Tallio Tissue)		Ť	0.54	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
355	736014					
		15 16 (Normal Calon vs. Calon T. T.				
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	6.00	5.68
356	646577	10,17 (Coton rumor rissue vs. Coton Metastasis)	6	0	6.09	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
357	732254					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	8		7.92
358	7037					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	9		9.76
359	7037					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	9		9.76
360	6937					
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	
361	7572					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.67
362	388085					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9.51	
363	2676					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	17		9.22
364	639240					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9.51	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
365	650472					
		1517.01			6.44	-
266	727700	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
366	727789					
		16 17 (Color Turner Tiesus us Color Material)			6.00	<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	6.09	5.68
367	2495	13,10 (Normal Colon vs. Colon Tumor Tissue)		-		3.08
307	2493			<u> </u>		<b>-</b>
		03,04 (Breast, High Met vs. Breast, Non-Met)	27	5	5.27	<u> </u>
		08,09 (Lung, High Met vs. Lung, Low Met)	13	2	9.08	<u> </u>
368	732254	l l l l l l l l l l l l l l l l l l l				
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	8		7.92
369	5268					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	15		16.26
370	11881					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
371	448677					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	1	11.63	
372	1876					
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	3	4.88	
373	3441					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		01,02 (Colon, High Met vs. Colon, Low Met)	4	13		3.52
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	0	7.81	
374	726134					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8	0.10	7.57
27.5	20.40	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
375	9048					ļ
		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	
376	26489	01,02 (Colon, Fright Met vs. Colon, Low Met)	+	Ů	0.10	
270	20102					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	8		8.2
377	644205					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
378	468689					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
379	638971				· · · · · · · · · · · · · · · · · · ·	
					0.50	
200	10074	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
380	10274					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
381	6725	05,04 (Breast, High Met Vs. Breast, Non-Met)	<del>-   •</del>			7.17
501	0.20					
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	
382	2488					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
383	8366			L		
		21,22 (Normal Prostate vs. Prostate Cancer) (Normal Prostate vs.	2	15		7.63
		Prostate Cancer) 03,04 (Breast, High Mct vs. Breast, Non-Met)	15	2	7.32	7.05
384	502683	05,04 (Bleast, High Met 75. Bleast, 11011 Met)	13		1.52	
			1			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
385	450914					
:		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
386	21205					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	9		9.22
387	644205					
		151601 101 7 7	+ -		7.4	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
200	5260	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
388	5268					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	15		16.26
		101,02 (Colon, 111gn Mice va. Colon, LOW Mice)		رب		10.20

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
389	8012					
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
390	11270					
			ŀ			
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
391	10924			<u> </u>		
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
393	3650			ļ		
						ļ
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	2	4.88	ļ
394	1655					ļ
		01,02 (Colon, High Met vs. Colon, Low Met)	67	2	30.9	<u> </u>
		21,22 (Normal Prostate vs. Prostate Cancer)	116	51	2.24	
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	58		59.45
395	3275					
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	0	7.81	
396	3355					
		03.04 (D	21	7	2.02	
207	2070	03,04 (Breast, High Met vs. Breast, Non-Met)	21	7	2.93	
397	2078		<del></del>			
		02.04 (Duncet Wish Making Duncet Non Mat)	11	2	5.37	
200	4800	03,04 (Breast, High Met vs. Breast, Non-Met)	11		3.31	
398	4809			-		<del>                                     </del>
		08,09 (Lung, High Met vs. Lung, Low Met)	3	27		6.44
		01,02 (Colon, High Met vs. Colon, Low Met)	15	1	13.84	0.44
399	6402	01,02 (Colon, 111gh Met Vs. Colon, Low Met)	- 13	<u> </u>	15.04	
377	0402					
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	
400	555244					<del></del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
401	548965					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14		3.45
402	4747					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	10		5.42
403	40208					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
404	14596					
		08,09 (Lung, High Met vs. Lung, Low Met)	14	6	3.26	
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	17		17.42
405	7110					
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	19		4.87
406	7110					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	19		4.87
407	6592					
		03,04 (Breast, High Met vs. Breast, Non-Met)	12	1	11.71	
408	6455	03,01 (Bloads, Tilgii Met 10. Bloads, 110ii Met)		Ė		
- 100	0400					
		102 04 (Preset High Motors Preset Non Met)	6	0	5.85	
400	2720	03,04 (Breast, High Met vs. Breast, Non-Met)	<del></del>	0	5.65	
409	2738			-		1
		OZ O	16	1	2.66	ļ
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	4	3.66	
410	696					
		03,04 (Breast, High Met vs. Breast, Non-Met)	37	13	2.78	ļ
		01,02 (Colon, High Met vs. Colon, Low Met)	24	87		3.93
411	379186					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
412	1588					
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	95		6.49
		01,02 (Colon, High Met vs. Colon, Low Met)	45	19	2.18	
413	7007					
	7007					·····
		01,02 (Colon, High Met vs. Colon, Low Met)	0	9		9.76
414	9025	01,02 (Colon, High Met 13. Colon, Low Met)				7.,,
414	9023					
		01 02 (Calan IViah Matana Calan Lam Mat)	7	0	6.46	-
44.7	(50540	01,02 (Colon, High Met vs. Colon, Low Met)		0	0.40	
415	650749					
				-	7.51	-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
416	553158					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
417	641703					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	0	12.88	
418	833					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	
		08,09 (Lung, High Met vs. Lung, Low Met)	18	9	2.79	
		03,04 (Breast, High Met vs. Breast, Non-Met)	16	3	5.2	
419	649259					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
420	451179					
.20	,			<del>                                     </del>		
		15.17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.38
421	0505	13,17 (11011) at Coloit 1155uc vs. Coloit Wiciastasis)		-		3.50
421	9505					
				1.		
		03,04 (Breast, High Met vs. Breast, Non-Met)	3	15		5.12
422	736728			ļ		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
	ļ	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.6
423	380412			1		
				1		
-		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	+
424	642425					1 -
						<b>-</b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	1
425	405073					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs Colon Tumor Tissue)	6	0	6.34	
426	174250				i	
						<b>T</b>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	13		6.4
427	726281				† ·	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	†
428	639029					<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
429	452245					<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	12	-	11.82
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	12		11.18
430	510254					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	17	1	17.97	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
431	642425					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
432	51939					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	28	3	9.87	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	28	0	30.05	
433	7379					
124	E46622	01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
434	546632					
-+		16 17 (C-1 T T				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	15	0	15.23	
435	734827	23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	3	34		11.22
433	134821		_			
		16.17.(C-1 T T' T' T' T'				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
126		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
436	2554				I	
		00.04/D				
127		03,04 (Breast, High Met vs. Breast, Non-Met)	20	7	2.79	
437	643285					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	

## Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
438	448770			T		
				1		1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<b></b>
439	375380					1
				$T^{-}$		1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8 38
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.86
440	726134			1		
				1		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
441	422687			† <del></del>		
				1	ļ	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.16	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	10		9.46
442	448436				<b> </b>	
			<del> </del> -	<del>                                     </del>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	20	2	10.16	<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	4	20		4.73
443	644893		<del></del>	<del>                                     </del>	<del> </del>	
				<del>                                     </del>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
444	559104			<del>  </del>		
	237101		+	<del>                                     </del>		<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
445	551172	15,75 (16,71m) Colon 15. Colon 14,1161 115546)		<del>اٽ</del>		3.00
	001112	<del> </del>	<del></del>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)		10		9.46
446	724296	10,10 (10) mar edon vs. edon ramor rissae)	<del></del>	<del>                                     </del>		2.40
	72.270	<del> </del>				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	128		121.1
<del></del>		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	128	0	129.99	121.1
447	735936	10317 (Colon Famor Fissac Vs. Colon McCastasis)	120	<u> </u>	129.99	
	733730					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1 9	0	9.14	0.51
448	556326	1917 (2010) 14/10/ 11/04/ 75/ 2010) 1/10/45/45/5	<del></del>	Ť	2.14	
	000000					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		8		7.88
449	729699	1017 (Colon Tanior Fisher Vs. Colon McCastasis)	<del></del>	۳		7.00
-17/	727077		<del></del>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)		6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	2.08
450	550694	10,17 (Colon Tuntor Tissue vs. Colon Pretastasis)	<del></del>	$\vdash$	0.09	
750	330074					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	21	7	3.22	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	21	1		
451	734738	15,10 (Normal Colon vs. Colon Tumor Hissue)	21		22.2	
401	134136			$\vdash$		
		16.17 (Colon Tumor Tioque va Colon Materia)			6.00	<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	5.70
453	404503	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
452	404502					
		151601 101 7		┝┋┩		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	

Table 5 SEQ **CLST** Library Pair A,B Á  $\overline{\mathbf{B}}$ A/B B/A 453 554151 15,17 (Normal Colon Tissue vs. Colon Metastasis) 1 15 13.97 15,16 (Normal Colon vs. Colon Tumor Tissue) 9 8.51 649852 454 15,17 (Normal Colon Tissue vs. Colon Metastasis) 6 0 6.44 455 734063 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 6.62 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 7 0 7.11 456 7279 0 01,02 (Colon, High Met vs. Colon, Low Met) 6 6.5 457 2676 01,02 (Colon, High Met vs. Colon, Low Met) 9.22 2 17 649148 458 0 8.59 15,17 (Normal Colon Tissue vs. Colon Metastasis) 8 459 1953 03,04 (Breast, High Met vs. Breast, Non-Met) 12.3 4 48 01,02 (Colon, High Met vs. Colon, Low Met) 39 13 2.77 460 650108 15,17 (Normal Colon Tissue vs. Colon Metastasis) 9 0 9.66 461 515350 15,16 (Normal Colon vs. Colon Tumor Tissue) 14 0 14.8 14 15,17 (Normal Colon Tissue vs. Colon Metastasis) 3 5.01 462 402494 15,16 (Normal Colon vs. Colon Tumor Tissue) 3 13 4.1 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 13 3 4.4 649148 463 15,17 (Normal Colon Tissue vs. Colon Metastasis) 0 8.59 8 464 833 03,04 (Breast, High Met vs. Breast, Non-Met) 16 3 08,09 (Lung, High Met vs. Lung, Low Met) 9 2.79 18 15,16 (Normal Colon vs. Colon Tumor Tissue) 12 3 4.23 465 139730 03,04 (Breast, High Met vs. Breast, Non-Met) 9.22 0 9 466 453079 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 1 8.12 8 467 546705 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 0 6 5.91 468 644903

Table	5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
469	732254					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.6
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	8		7.9
470	561180					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	
471	732254			L		
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	8		7.9
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.6
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	L
472	449204					<u> </u>
		<u> </u>				<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	14		13.
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	14		4.
473	185651					L
						L
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.1
474	639029					
						<b> </b> -
	150001	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<u> </u>
475	452986			<u> </u>		
		15160			0.46	
126	720770	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
476	729779					
		15 16 (Normal Calar va Calar Truman Tirana)	<del></del>	8		7.5
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	0	0.13	7.5
477	646248	10,17 (Colon Turnor Tissue vs. Colon Metastasis)	8	0	8.12	<u> </u>
4//	040248					├
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<del> </del>
478	650448	10,17 (Cololi Tullor Tissue vs. Cololi Metastasis)			0.09	
7/6	030440					<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
479	642049	TOTAL (TOTAL COOK TISSUE TO: COOK TISSUES)		<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
	<del></del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
480	728273					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.6
481	446139					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	0	13.74	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	12	0	13.71	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	0	13.95	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	12	0	10.26	
ſ						
482	2783		(			

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	5	40		8.2
		01,02 (Colon, High Met vs. Colon, Low Met)	27	6	4.15	
483	642906					
						1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
484	8332					
						J
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
485	453470					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	1	12.88	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	
486	552277			<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
487	464029					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
489	649722					
	·	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
490	612572					
						L
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
491	385980					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	27	12	2.28	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	27		12.77
492	141185					L
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	L
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	L
493	463824					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
494	446139				<del></del>	<u> </u>
					12.54	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	0	13.74	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	0	13.95	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	12	0	13.71	<u> </u>
105	725004	18,20 (Normal Colon Tissue vs. Colon Metastasis)	12	0	10.26	
495	725994					
		15 16 Olympia City To The City				5.60
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	6.00	5.68
406	726670	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
496	736679	<del> </del>				
		16 17 (Colon Tumor Tioque va Colon Material)			6.00	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	6	6.09	5 (0
497	551718	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
498	640525					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
499	645210					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	1	10.57	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
500	6567					
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	
501	646146					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
502	4934					
		01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76
503	450791					
		<del> </del>				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	18	6	3.17	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	18	3	6.44	
504	227936					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
505	9436					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	15		3.69
506	2557					
		01,02 (Colon, High Met vs. Colon, Low Met)	23	8	2.65	
507	11356					
						<del>                                     </del>
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
508	7571					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.67
509	558116					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5 91
510	216574					
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	15	2	6.41	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	107	29	3.96	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	107	29	3.9	
511	455145					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
512	649148	, , , , , , , , , , , , , , , , , , ,	<del> -</del> -	<u>*  </u>	3.17	
<del></del>		<del> </del>		$\vdash \vdash \vdash$		
ı		<del> </del>				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0 1	8.59	

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Table 5						
SEQ	CLST	, , , , , , , , , , , , , , , , , , ,	A	В	A/B	B/A
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
<del></del>	<del>                                     </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
514	304253					
	<u> </u>					
	640747	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
515	649717			_		
	<del> </del>	1/12/01 E				
516	5020	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
516	5838			<del> </del>		<b>_</b>
	<u> </u>	01 02 (Colon Hick Mayor Colon I. May				
517	454050	01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76
317	434030			┿	ļ	<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		+-	0.50	<del>}</del>
518	557903	13,17 (Normal Colon Tissue vs. Colon ivietastasis)	8	1	8.59	
J10	337703					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	<del>-   _</del>	-	<del>                                     </del>	0.61
519	1724	13,10 (1011Mai Colon Vs. Colon Tullor Tissue)	0	9		8.51
	1/21			+		
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	6	24	<u> </u>	4.1
520	734803	(2. table) 21g. 11et 10. Stouds, 1ton 11ety		1 24	<del> </del>	4.1
				<del></del>		-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.07	5.68
521	557948			<u> </u>		3.00
			<u> </u>	+-		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
522	5838					
		01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76
523	2334					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
524	450953					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	14		6.52
	1010	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	14		13.79
525	4840					
		02.04 (P				
526	728421	03,04 (Breast, High Met vs. Breast, Non-Met)	18	6	2.93	
220	120421					
		15,16 (Normal Colon vs. Colon Tumor Tissue)				F. (C.
<del></del>	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	6.00	5.68
527	4747	10,17 (Colon Tullior Tissac vs. Colon Metastasis)	6	0	6.09	
	,					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	10		5.42
528	648934	, (,g.,to, Colon, Lon Met)		10		3.42
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
529	1787			<u> </u>		
					<del></del>	
		01,02 (Colon, High Met vs. Colon, Low Met)	17	36		2.3

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SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
530	558098				<u> </u>	
				$\dagger$		+
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
531	1655			T		_
				1	<b>†</b> – –	
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	58	<del>                                     </del>	59.4
		21,22 (Normal Prostate vs. Prostate Cancer)	116	51	2.24	1
		01,02 (Colon, High Met vs. Colon, Low Met)	67	2	30.9	+-
532	158601			+	1	
				1	<del></del>	† –
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	10	<del>                                     </del>	10.2
533	185486			+	<del>                                     </del>	1
					<del> </del>	+
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	13		13.3
534	7110			<del>                                     </del>	<del>                                     </del>	13.3
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	19		4.87
535	2543					1.07
				<u> </u>	<u> </u>	+ -
		01,02 (Colon, High Met vs. Colon, Low Met)	23	7	3.03	+
536	115762					
				<u> </u>	<del>                                     </del>	
		21,22 (Normal Prostate vs. Prostate Cancer)	16	5	3.15	<del>                                     </del>
*		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
537	696					0.52
		01,02 (Colon, High Met vs. Colon, Low Met)	24	87		3.93
		03,04 (Breast, High Met vs. Breast, Non-Met)	37	13	2.78	
538	1948					
		03,04 (Breast, High Met vs. Breast, Non-Met)	37	15	2.41	-
539	696					<u> </u>
		01,02 (Colon, High Met vs. Colon, Low Met)	24	87		3.93
		03,04 (Breast, High Met vs. Breast, Non-Met)	37	13	2.78	
540	696					
		03,04 (Breast, High Met vs. Breast, Non-Met)	37	13	2.78	
		01,02 (Colon, High Met vs. Colon, Low Met)	24	87		3.93
541	380477					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
542	638799					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
543	551982					
		6,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
544	551982			$\neg$	<del></del>	
			<del>-    </del>	_		
		6,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	<del>  </del>	5.91
545	521840			-		2.71
	T			_		

Table 5

Table 5	5					
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
546	561180					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	
547	556245			1		<del> </del>
				† – –		
<u> </u>		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	<del> </del>	5.68
548	449792	Total (Total Cook to Cook Taking Tibbe)	Ť	<del>                                     </del>		2.00
3.0	112722			+ -	<del> </del>	<del>                                     </del>
<u> </u>		19,20 (Colon Tumor Tissue vs. Colon Metastasis)		5	<del> </del> -	6.68
}		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	13		4.1
549	549722	13,10 (Normal Colon vs. Colon Tumor Tissue)	<del> -</del> -	113	<del> </del>	7.1
347	349122			-		<del> </del> -
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		6	<del> </del>	5.91
550	612572	10,17 (Coloit Tuttor Tissue vs. Coloit Metastasis)		10	<del> </del>	3.91
330	012372			<del>├</del> ──	<del> </del>	<del> </del>
		16 16 Oleman Colon Colon Transactions	<del></del>	<del> </del>	(24	<del> </del>
551	551225	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<del> </del>
551	551235		<del></del>	├	<u> </u>	<del> </del>
		16 16 (G 1 - T - T)	<del></del>	<del> </del>		600
552	440701	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7	<del> </del>	6.89
552	449701			├	ļ	<u> </u>
		16 17 (Color Torres Tirres Color Metadoria)	17	<del>├.</del> -	17.26	
<del> </del>		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	17	1	17.26	16.00
F52	275200	15,16 (Normal Colon vs. Colon Tumor Tissue)	1	17		16.08
553	375380		<del></del>	├	<u> </u>	<del> </del>
		16.12 (Color Town Times Color Materials)		<u> </u>	<b></b>	9.96
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9	<del> </del>	8.86
554	56940	15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.38
334	30940		<del></del>	<del> </del>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)			<del> </del>	5.60
				6	<u> </u>	5.68
555	549160	15,17 (Normal Colon Tissue vs. Colon Metastasis)	- 0	8		7 45
333	349100			-		
		15 17 Ob 1 C-1 Ti C-1 M destrois				( 52
554	554151	15,17 (Normal Colon Tissue vs. Colon Metastasis)	- 0	7		6.52
556	334131					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		1.5		12.07
<del> </del>			1	15 9		13.97
557	727221	15,16 (Normal Colon vs. Colon Tumor Tissue)	<del></del>	9		8.51
557	727331			$\vdash$		<del> </del> -
<del>  </del>		16.17 (Colon Turnor Tirror or Colon Maria	<del> </del> -	<u> </u>	( 00	<del> </del>
<b> </b>		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	5.00
550	661600	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
558	551502			$\vdash \vdash$		<del>                                     </del>
$\vdash$		15 14 Olympia Colon va Colon Town Tillian		-	2.22	<del> </del>
├──┤		15,16 (Normal Colon vs. Colon Tumor Tissue)	22	7	3.32	0.50
F50	612572	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	19		2.67
559	612572			<b> </b>		<u> </u>
<del>  </del>						
7.5	70:22:	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
560	701221			$\vdash$		
		16.17.60.1				
	270011	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	<u> </u>
561	378041					

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Table 5

SEQ	CLST	Library Pair A,B	A	B	A/B	<b>B</b> /A
	<del> </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		+		—
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	-	3.3	12
562	503491	(10711 Colon vs. Colon Turnor 113suc)		13		12.
				_		-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	1
563	452833					
561	640074	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
564	640974				ļ	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)				<b> </b>
565	735326	13,17 (Normal Coloit Tissue vs. Coloit Metastasis)	9	10	9.66	-
				-		-
		15,16 (Normal Colon vs. Colon Tumor Tissue)		7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	0.02
566	555944			T-		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
567	447532					
		15.16.0V1.0.1			ļ	
		15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.63	<u> </u>
568	455598	15,17 (Normal Colon Tissue vs. Colon Metastasis)		1	11.81	
	100070			-	<u> </u>	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	<del> </del>	5.91
569	555734		<del>-   •</del>	+ <u> </u>		3.51
				+-	<u> </u>	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	10		9.32
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	10		9.85
570	446663					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)				
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	6	32		5.25
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		9.36
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	32		2.13
571	449862			1-		2.13
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
572	549591					
		16 17 (C-1 T				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	24		3.38
573	553877	15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	24		11.18
				1—1		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
574	553501		<del>-   `</del>			1.00
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	1	14.8	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	3	5.01	
575	1905					
		01.02 (Color High May				
		01,02 (Colon, High Met vs. Colon, Low Met)	7	21		3.25
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52

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Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
576	446599					
					T T	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	2	6.87	
577	559409					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
578	551982			<u> </u>		<b>†</b>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
579	559057					
****				1		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
580	446760			<del>                                     </del>		0.52
				<del>                                     </del>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	22	7	3.19	
581	551502			╁╧	3.17	-
				<del> </del>	<u> </u>	<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	19		2.67
-		15,16 (Normal Colon vs. Colon Tumor Tissue)	22	7	3.32	2.07
582	446531	, ( and a second residue)		<del>                                     </del>	3.32	_
				<del> </del>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
583	506744	( and the state of	<del>-   `</del>	-		3.91
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8	7.4	7.88
584	401849	, ( )	Ť	-		7.88
	-	15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	15	7.4	14.77
585	453848	2-19-7 (Coton Variet 115050 15)		13		14.77
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	14		6.62
586	456764	to the state of th		14		0.02
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	1	15.03	·
587	446371		- 14	-1-	13.03	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
588	406413	20,10 (110 man colon 10. Colon Tumor Hasue)		-	0.34	
				<del> </del>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	-   1	9		8.86
589	555103	15542 VS. Colon Victasiasis)		-		8.80
-	000100					
<del>-  </del>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	<del>                                    </del>	7		6.52
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	-+		6.52
590	735292	-5,5. (Avortinal Eding Tissue vs. Eding Luthol Tissue)	0	6		5.94
370	133434					
		15 16 (Normal Colon vs. Colon T. T.				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
591		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
391	558534					
		1610/0				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89

639662

607

Table 5 SEQ CLST Library Pair A,B  $\mathbf{A}$ В A/B B/A 727181 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 6 0 6 09 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 5.68 593 551117 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 0 6 5.91 594 464040 15,16 (Normal Colon vs. Colon Tumor Tissue) 6 0 6.34 595 446371 15,16 (Normal Colon vs. Colon Tumor Tissue) 6 0 6.34 596 728408 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 12 0 12.19 15,16 (Normal Colon vs. Colon Tumor Tissue) 0 12 11.35 597 649259 15,17 (Normal Colon Tissue vs. Colon Metastasis) 7 0 7.51 15,16 (Normal Colon vs. Colon Tumor Tissue) 7 0 7.4 598 15414 15,17 (Normal Colon Tissue vs. Colon Metastasis) 9 8.38 599 639240 15,17 (Normal Colon Tissue vs. Colon Metastasis) 0 9.66 15,16 (Normal Colon vs. Colon Tumor Tissue) 9 0 9.51 549722 600 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 0 6 5.91 601 561499 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 8 1 7.88 15,16 (Normal Colon vs. Colon Tumor Tissue) 8.46 602 639029 15,17 (Normal Colon Tissue vs. Colon Metastasis) 6 0 6.44 449512 603 15,17 (Normal Colon Tissue vs. Colon Metastasis) 4 14 3.26 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 1 14 13.79 446987 604 15,17 (Normal Colon Tissue vs. Colon Metastasis) 10.73 10 0 605 466302 15,16 (Normal Colon vs. Colon Tumor Tissue) 9 9.51 553802 606 15,17 (Normal Colon Tissue vs. Colon Metastasis) 1 9 8.38

7.51

15,17 (Normal Colon Tissue vs. Colon Metastasis)

Table 5

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Table 5						
SEQ	CLST		A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
608	551527			ļ		ļ
		15 17 Olement Calon Times of Calon Material				
609	730389	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7	<u> </u>	6.52
-007	750507			╂	<u> </u>	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9	7.14	8.51
610	640974			<u> </u>		0.51
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
611	417155					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
612	417155					
		15 16 Oleman Color Color Transport		<u> </u>		
613	451784	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
015	431764					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.90
614	649152	(Count I amor 110000 to Coton Madasasis)				6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
615	450867					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
616	143436					
		15 17 O				
617	549395	15,17 (Normal Colon Tissue vs. Colon Metastasis)	21	9	2.5	
017	349393					
	-	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	13		12.8
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	13		6.06
618	639273			- 13		0.00
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9 51	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
619	506744					
		15 16 (Normal Calan vs. Calan T				
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	74	7.00
620	736595	10,17 (COLOR TURNOT TISSUE VS. COLOR IVICIASIASIS)	0	8		7.88
				-		
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	5.57	5.68
621	230995					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
622	451784					
622		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
623	226324					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	<del>-   _  </del>			
		- v, . , (Colon Turnor Traduc vs. Colon Miciastasis)	0	6		5.91

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	
624	449617			1		T
						T
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	18		1
625	451092					T
						1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		
626	546642					
						L
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	<u> </u>
627	553736					┡
		16.17 (Color Times Tierra of Color Metadois)		10		-
628	394413	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	10		<u> </u>
028	394413			-		-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	113	3	39.81	-
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	21	0	17.95	┢╌
		18,19 (Normal Colon Tissue vs. Colon Tumor)	21	2	12	$\vdash$
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	113	0	121.29	Π
629	556326					Г
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7
630	448606					<u> </u>
						_
(2)	2011	15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	20		6
631	394413					_
	<del></del>	15 17 (Normal Colon Ticque us. Colon Materialia)	112		131.30	$\vdash$
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 18,19 (Normal Colon Tissue vs. Colon Turnor)	21	2	121.29	├
		15,16 (Normal Colon vs. Colon Tumor Tissue)	113	3	39.81	┝
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	21	0	17.95	
632	645633	( ( ( ( ( ( ( ( ( ( ( ( ( ( ( ( ( ( (		- Ŭ	17.55	$\vdash$
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
633	551634					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5
634	556326					
625	£40707	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7
635	540787				···-	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	ļ
+		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.4	
636	648872	25,1, (170111111 COIOII 113511C 13. COIOII IVICIASIASIS)			1.31	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
637	643804					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
638	446139					
			I	- 1		

Table 5 Page 41 of 155

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	12	0	13.71	T
		18,20 (Normal Colon Tissue vs Colon Metastasis)	12	0	10.26	$\top$
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	0	13.74	T
639	640356					╁
	ļ					
(10	270197	15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	$oldsymbol{\downarrow}$
640	379186					╀
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	╁
641	454927				0.02	
		1618 (61 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7				L
642	401849	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	14	<u> </u>	╀
	101015					╁
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	15		1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
643	452414			ļ		$oldsymbol{\perp}$
		15,16 (Normal Colon vs. Colon Tumor Tissue)	17	0	17.97	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	17	0	18.25	$\vdash$
644	446789	Total Colon Tiber (i. Colon Medians)	17	<u> </u>	16.23	╁
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	5	3.38	L
645	189561					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9	** • **	H
		08,09 (Lung, High Met vs. Lung, Low Met)	1	14		1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9		1
646	640323					
		15 17 01 10 1				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	L
647	558116	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	H
						├
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		-
648	468109					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)			6.44	<u> </u>
649	481441	10,11 (Normal Colon 11880e vs. Colon Metastasis)	6	0	6.44	$\vdash$
						$\vdash$
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	12	0	12.19	
650	449956					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	_
651	727224		10		10.10	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
652	551007	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7
652	551907					
-+		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
653	447532					
ſ		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	0	11.63	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	]
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	1	11.81	
654	447532					
						上
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	1	11.81	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	0	11.63	_
655	558454					_
		48.5.404.5.404.		<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		<del>  '</del>
656	502683			_		╂_
		151(0) 101 01 5				┼
657	446000	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	╁
657	446909			-		╀
	······································	16 17 (Color Transa Tiransa Color Material)		12		Η,
658	452506	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	12	<u> </u>	-
036	432300					╁
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	╁
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5
659	449792	1.55 (Colon Tulior Tiodae 15. Colon Intelastacia)	-   '	۳		Η.
						$\vdash$
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	0	5		-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	13		
660	549395					T
						T
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	13		6
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	13		1
661	234653					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	18		2
662	453911					L
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5
663	452071					<u> </u>
						ļ
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	14	0	11.97	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	109	1	115.21	
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 18,19 (Normal Colon Tissue vs. Colon Tumor)	109	0	117	<del> </del>
664	451032	10,17 (NOTHAL COLOH TISSUE VS. COLOH TUMOT)	14	0	16	$\vdash$
004	431032					$\vdash$
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
665	446680	17,10 (10) that Colon 15, Colon Tullor 11550C)		<u> </u>	U.J-1	$\vdash$
			<del></del>			$\vdash$
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	29	84		1
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	40	94		2
666	641884					<u>-</u> -
						$\vdash$
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	Г
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
667	452800					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	13		4
668	461835					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
669	548965					Т
-		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14		
670	734793					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	_
671	539955					╙
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	3	47		1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	24	0	24.37	+ 1
		15,17 (Colon Turior Tissue vs. Colon Metastasis)		+		╁
672	561892	13,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	╄
072	301092			-		╀
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6
673	562292					Ť
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5
674	420686			ļ		_
		16 17 (Calar Tarras Tirras Calar Maria		<u> </u>		<u> </u>
675	9436	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7
0/3	9430					1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	15		3
676	1013			15		H
						$\vdash$
		01,02 (Colon, High Met vs. Colon, Low Met)	40	84		2
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	Г
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
677	412364					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	L
678	44424	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	_
0/8	44424					
		08,09 (Lung, High Met vs. Lung, Low Met)	4	40		7
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	25	90		3.
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	41	90		2.
679	394413					Ť
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	113	0	121.29	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	21	2	12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	113	3	39.81	
(00	440617	18,20 (Normal Colon Tissue vs. Colon Metastasis)	21	0	17.95	
680	449617					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	10		2
681	455032	10,17 (Colon Tumor 1155ue vs. Colon Metastasis)		18		2.
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.
682	185400					
- 1		03,04 (Breast, High Met vs. Breast, Non-Met)	1	62		63

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
683	453911					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
684	650297					
	<u> </u>					
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
685	185400					
	<u> </u>					1
	ļ	03,04 (Breast, High Met vs. Breast, Non-Met)	1	62		63.55
(0)	440512	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	1	9		8.86
686	449512			-		<del>                                     </del>
		15 17 01 1 C 1 T		+		<b>-</b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14	<u> </u>	3.26
687	44424	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	1	14	<u> </u>	13.79
	44424			<del>                                     </del>		ļ
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	41	90		2.16
· · · · · · · · · · · · · · · · · · ·		15,17 (Normal Colon Tissue vs. Colon Metastasis)	25	90		3.35
		08,09 (Lung, High Met vs. Lung, Low Met)	4	40		7.16
688	556216	(		1		7.10
				1		l
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
689	448677					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	1	11.63	
690	375380					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.38
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.86
691	379341					
				ļ		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	14		4.35
602	276000	08,09 (Lung, High Met vs. Lung, Low Met)	2	21		7.51
692	376988					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	<del></del>			7.00
693	559806	10,17 (Coloil Tulliol Tissue vs. Coloil Metastasis)	1	8		7.88
0,0	227000					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
694	550195	· · · · · · · · · · · · · · · · · · ·		<u> </u>		3.71
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
695	562221					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
696	211					
		01,02 (Colon, High Met vs. Colon, Low Met)	109	206		2.05
		03,04 (Breast, High Met vs. Breast, Non-Met)	121	43	2.75	
697	6751	l	1	, 1		

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	11		11.27
		08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	
698	6751					
	<u> </u>	08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	
				11	0.99	11.27
700	2883	03,04 (Breast, High Met vs. Breast, Non-Met)	- V	11		11.2
						t
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	21		2.39
701	9784					
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
702	649722					
		16170			0.50	<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.59 6.09	
704	10340	1017 (COOR TURIO TISSUE 13. COOR MCLUSIUSIS)		Ů	0.07	
705	1640	03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
705	1649					<del> </del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	1	14.63	
706	4325					
		02 04 (December 1912) Marcon December 1912	7		( 92	
707	10882	03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
	10002					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	20		3.42
708	10342					
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
709	6474					
710	10240	01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
710	10340					$\vdash$
·		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
711	734723					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9	9.14	8.51
712	452142					
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	10	5.00	9.46
713	185432	16,17 (Colon Turnor Hissue vs. Colon Metastasis)	10	2	5.08	
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	26		26.65
714	11456					<u> </u>
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
715	508892	, , , , , , , , , , , , , , , , , , ,				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	10		4.92
716	67					
		03,04 (Breast, High Met vs. Breast, Non-Met)	23	0	22.44	
		01,02 (Colon, High Met vs. Colon, Low Met)	0	32		34.6
717	2636					
		08,09 (Lung, High Met vs. Lung, Low Met)	7	1	9.78	
		01,02 (Colon, High Met vs. Colon, Low Met)	8	19		2.57
718	735028					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.6
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
719	1924					
		01,02 (Colon, High Met vs. Colon, Low Met)	8	21		2.8.
720	640116					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	4	5.02	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	19	0	20.39	
721	6546					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	9		9.7
722	730866					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.5
723	4829					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.5
724	546632					
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	3	34		11.2
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	15	0	15.23	Ì
725	549934					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3.88	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	20		6.5
		21,22 (Normal Prostate vs. Prostate Cancer)	8	0	7.87	
726	649655					
			ŀ			
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	12	0	12.19	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
727	62016					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
728	2783					
	,	01,02 (Colon, High Met vs. Colon, Low Met)	27	6	4.15	
		03,04 (Breast, High Met vs. Breast, Non-Met)	5	40		8.2
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
729	3876		<u> </u>			-
			<del></del>			
<del></del>		21,22 (Normal Prostate vs. Prostate Cancer)	9	26		2.9

Table 5

CLST	Library Pair A,B	A	В	A/B	B/A
	15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	14		4.35
20036					
	151501 101 7		ļ.,		
644022	15,17 (Normal Colon Tissue vs. Colon Metastasis)		13		12.11
044032			+-		<del> </del>
	16.17 (Colon Tumor Tissue vs. Colon Metastasis)	124	0	125 92	
			+	123.72	16.76
	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
451636					
	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
3428			ļ		<u> </u>
	02 04 (Prooct High Motors Procest Nov. Mas)	20	-	10.51	-
643054	03,04 (Breast, High Met Vs. Breast, Non-Met)	- 20	0	19.51	<u> </u>
043334					
	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
456506					<b> </b>
	15,17 (Normal Colon Tissue vs. Colon Metastasis)	17	7	2.61	
449269					
		26		3.99	1
723713	15,16 (Normal Colon vs. Colon Tumor Tissue)	26	5	5 5	
132/12					-
	15.16 (Normal Colon vs. Colon Tumor Tissue)	- 0	7		6.62
		7	-	7.11	0.02
696					
	01,02 (Colon, High Met vs. Colon, Low Met)	24	87		3.93
	03,04 (Breast, High Met vs. Breast, Non-Met)	37	13	2.78	
456528					ļ
	15 16 Olympia Color To Tr		- 1		
4043	15,16 (Normal Colon Vs. Colon Tumor Tissue)	0	'		6.62
	01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76
3639					
	03,04 (Breast, High Met vs. Breast, Non-Met)	12	3	3.9	
1024					
					L
			<del></del>	£ 21	299.28
1247	U6,U9 (Lung, High Met vs. Lung, Low Met)	41	11	5.21	
1241					
	03,04 (Breast, High Met vs. Breast, Non-Met)	51	15	3.32	
4934	Total Transport Control of the Property of the			5.52	
	01.02/01 17.13/4 01 1 3/4				0.77
	01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76
	20036 644032 451636 3428 643954 456506 449269 732712 696 456528 4043 3639 1024	15,17 (Normal Colon Tissue vs. Colon Metastasis)	15,17 (Normal Colon Tissue vs. Colon Metastasis)   3	15,17 (Normal Colon Tissue vs. Colon Metastasis)   3   14	15,17 (Normal Colon Tissue vs. Colon Metastasis)   3   14

Tal	nle	5

Table 5	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLSI	03,04 (Breast, High Met vs. Breast, Non-Met)	56	5	10.93	2/12
746	452726	05,04 (Bleast, High Met Vs. Bleast, Mon-Met)				
740	432120					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	
747	725825					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
748	456808					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	42		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	42	1	42.65	
749	729295					
						( (2
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	7.11	6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
750	551907			-		
		15 16 Olympia Colon va Colon Turnor Tiggue)	7	0	7.4	
751	551527	15,16 (Normal Colon vs. Colon Tumor Tissue)		<del>- </del>	7	
751	551527					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
752	7098	13,17 (Normal Colon Fissue vs. Colon Metadado)				
132	7038					
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	
753	4589					
,,,,,						
		01,02 (Colon, High Met vs. Colon, Low Met)	14	2	6.46	
754	554812					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
755	3114					
				<u> </u>	0.70	
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	ļ
756	6031			<del> </del>		ļ
		101 02 (C. L. Hick Maters Colon LevelMat)	9	1	8.3	
757	195629	01,02 (Colon, High Met vs. Colon, Low Met)		<u> </u>	0.5	
757	185628			<u> </u>		<u> </u>
	<del>                                     </del>	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
758	24719	25,5 . (21000), 11811 1151 1151 21500), 1001 1150)		1		
,,,,,				<u> </u>		
	<b> </b>	03,04 (Breast, High Met vs. Breast, Non-Met)	4	26		6.66
	1	21,22 (Normal Prostate vs. Prostate Cancer)	4	14		3.56
759	3428					
		03,04 (Breast, High Met vs. Breast, Non-Met)	20	0	19 51	
760	2676					
				1		<u> </u>
		01,02 (Colon, High Met vs. Colon, Low Met)	2	17		9.22
761	649148			<del> </del>		<u> </u>
<u> </u>	<b></b>			<u> </u>	2.52	<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	<u> </u>

The first of the f

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
762	234605					
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	17	5	3.43	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	39	14	2.83	-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	32	14	2.45	
5.60	2224	13,17 (Normal Colon Tissue vs. Colon Freuestation				
763	2224					
		CO O4 (D 4 IV-l- Mat up Broost Non Met)	44	8	5.37	
		03,04 (Breast, High Met vs. Breast, Non-Met)		<del>-  </del>		
764	185642					<del></del>
		D . N . MA	0	7	-	7.17
		03,04 (Breast, High Met vs. Breast, Non-Met)	- 0			7.17
765	649655					
					6.44	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	12	0	12.19	
766	2854					
		03,04 (Breast, High Met vs. Breast, Non-Met)	31	3	10.08	-
767	453470					<u> </u>
						ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	1	12.88	
768	11012					
	1					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
769	535208	01,02 (Coton, 11g. 11)				
709	333208					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
770	119606	13,17 (Normal Colon Fissac vs. Colon Fictasacis)				
770	448606					
		15 17 Olympia Colon Metactoric)	3	20		6.21
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		20		<del>                                     </del>
771	12304					$\dagger$
		The state of the s	5	17		3.48
		03,04 (Breast, High Met vs. Breast, Non-Met)		1,	<del></del>	+
772	2756			-		+
	<u> </u>			0	2 22	+
		01,02 (Colon, High Met vs. Colon, Low Met)	28	8	3.23	+
773	367			-	<del> </del>	+-
	<u> </u>				<u> </u>	1 2 2
		08,09 (Lung, High Met vs. Lung, Low Met)	30	99	127	2.3
		03,04 (Breast, High Met vs. Breast, Non-Met)	105	24	4.27	+
		01,02 (Colon, High Met vs. Colon, Low Met)	19	97		5.5
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	4	3.44	<del> </del>
774	11351					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
775	6858					
.,,	- 5550					
	+	01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.6
776	7750	organ (Colon) regin rate to: Colony 251 1225)				
//0	1/30			1	1	1
	-	03,04 (Breast, High Met vs. Breast, Non-Met)	1	14	<del>                                     </del>	14.
	1	105,04 (Dicast, night wife vs. Dicast, Non-Mich	1 -	1		-

Table 5

SEQ	CLST	Library Pair A,B	A	B	A/B	B/A
		02.04 (December 19:1.14)		<u> </u>		ļ <u>.</u>
778	11552	03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	<b>!</b>
778	11332			<del> </del>		-
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
779	12448					
	Ì					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
781	10342					
	ļ					
702	0026	03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6 83	
782	9026			-		
		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	<u> </u>
783	10342	01,02 (Colon, Tright Wet Vs. Colon, Low Wet)	<del></del>	0	0.40	
, , ,	100.2		,			
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
784	6455					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
785	6455					
786	2/16	03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
760	3416				<u></u>	
		01,02 (Colon, High Met vs. Colon, Low Met)	1	11		11.93
787	3416	on, or (colon, high lifet to colon, how lively				11.93
		01,02 (Colon, High Met vs. Colon, Low Met)	1	11		11.93
788	2889					
				<u> </u>		
700	7202	01,02 (Colon, High Met vs. Colon, Low Met)	8	19		2.57
789	7393					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		0.2
790	14390	05,04 (Bicast, High Mct vs. Bicast, Non-Met)		°		8.2
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	43		22.04
791	661					
		03,04 (Breast, High Met vs. Breast, Non-Met)	77	10	7.51	
703	452000	08,09 (Lung, High Met vs. Lung, Low Met)	0	10		7.16
792	452992					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7 51	
793	1943	15,17 (Monthal Colon 11550C vs. Colon Mclastasis)		0	/ 31	
						<del>-</del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	16	4	3.9	
		01,02 (Colon, High Met vs. Colon, Low Met)	9	29		3.49
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	19		3
	2027					
794						
794		03,04 (Breast, High Met vs. Breast, Non-Met)	12	35		

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
				1	1	1 2/11
		01,02 (Colon, High Met vs. Colon, Low Met)	1	8		8.67
796	650493					
		15 17 01 10 1 7		_	ļ	
797	640318	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
- <i>''</i>	040318			-	<del>                                     </del>	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<del> </del>
798	646309	( Colon Made Colon )		<del>                                     </del>	0.44	<u> </u>
				$\dagger$		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
799	4316					
800	140701	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
800	449701			_	ļ	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		<del> </del>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	17	1	17.26	16.00
801	560367	13,10 (Normal Colon Vs. Colon Fullor Tissue)	1	17	<del> </del>	16.08
				+		ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9	<u> </u>	8.51
802	9997			1		
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
803	649106					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)				
804	461835	13,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
				-		
		15,16 (Normal Colon vs Colon Tumor Tissue)	7	0	7.4	
805	640590			i		
	<del></del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
906	648340	15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
806	048340					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
807	554812	Today 13. Colon Medicastasis)	-   0	'	0.44	
				<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
808	447035					
		16.17.(O.)T. T'				
809	1208	16,17 (Colon Tumor Tissue vs. Colon Metastasıs)	8	1	8.12	
557	1200		-	<u> </u>		
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	0	12.68	
810	3114	(2) (2) (2) (3) (4) (4) (4) (4) (4) (4) (4) (4) (4) (4	13		12.00	
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
811	3114					
912		03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
812	734078					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		1516 OL 1G1 G1 T		<u> </u>	<u> </u>	
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.62
813	450323	10,17 (Colon Turnor Tissue vs. Colon Metastasis)	7	10	7.11	<del> </del>
	1			-		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8	<del>                                     </del>	7.45
814	11567		,	Ť		7.43
				1		
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
815	11567					
916	((())	03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	ļ
816	6660					<u> </u>
		01,02 (Colon, High Met vs. Colon, Low Met)		<u> </u>	6.46	ļ
817	9026	1,02 (Colon, High Met vs. Colon, Low Wet)	7	0	6.46	-
				-	<u> </u>	
		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	
818	185539					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	10		10.25
819	3224					
		01.02.62.1 IX 1.34 G.1 X				
820	95700	01,02 (Colon, High Met vs. Colon, Low Met)	17	2	7.84	
020	93700			<del> </del>	ļ	
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	21		21.52
821	4439	, ( , , , , , , , , , , , , , , , , , ,		21		21.32
			<del> </del>			
		08,09 (Lung, High Met vs. Lung, Low Met)	10	2	6.99	
		01,02 (Colon, High Met vs. Colon, Low Met)	15	2	6.92	
822	3428					
		02.04 (D				
823	1456	03,04 (Breast, High Met vs. Breast, Non-Met)	20	0	19.51	
	1430					
		01,02 (Colon, High Met vs. Colon, Low Met)	9	22		2.65
		03,04 (Breast, High Met vs. Breast, Non-Met)	50	9	5.42	2.03
824	11343		30		3.42	
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
825	729206					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	10		9.46
826	558371	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.16	
020	336371					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	·
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
827	451589	A A A A A A A A A A A A A A A A A A A	10		3.37	
		15.17 (Normal Colon Tionus en Colon Material 1)	<del> </del>			
828	404475	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7	i	6.52

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SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
~- <del>C</del>		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	2	5.59	1 27.11
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	19	2	10.2	<del>                                     </del>
829	734582	15121 (1.10111111 (1.10111111111111111111111		<del>  -</del>		<del>                                     </del>
				1	<del>                                     </del>	<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	T
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
830	729779					
						ļ — —
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
831	555244					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
832	449269					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	26	7	3.99	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	26	5	5.5	
833	4609					
	<u> </u>				Ļ	
		01,02 (Colon, High Met vs. Colon, Low Met)	2	12		6.5
834	640318					
				<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
835	729851			<u> </u>	ļ	
	<del> </del>		<del></del>	<u> </u>		<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
026	11020	15,16 (Normal Colon vs. Colon Tumor Tissue)	- 0	7		6.62
836	11028		<del></del>			
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
837	643924	01,02 (Colon, High Met Vs. Colon, Low Met)	<del></del>	-		0.5
657	043924			<del> </del> -	<u> </u>	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<del></del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
838	630259		<del>+-</del>			<del></del>
				ļ —		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
839	11286				<del></del>	
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
840	185651					
	1	03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
			1			
841	7379					
841	7379					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
841	7379	01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	12		6.5
842	728408				12.19	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	12	12.19	
842	728408	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	12	12.19	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	_
	-	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	+
845	185489		+	╁	0.44	+
				1-		+
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	12		╅
846	447326					
847	11006	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	_
041	11006			-	ļ	+
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		╬
848	6863	, (,,,,,,,	-   <u>°</u>	+ •	<del>                                      </del>	╁
				$\vdash$		+
		01,02 (Colon, High Met vs. Colon, Low Met)	1	8		+
849	11351					
850	401553	01,02 (Colon, High Met vs. Colon, Low Met)	0	6		—
830	401333			<u> </u>		+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	10	8.59	┿
851	504513	, ( Section Metabolists)	+ -	<del>L</del> –	6.39	╁
				<u> </u>		+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	T
852	645979					
		1/17/01 7		<u> </u>		$\perp$
853	6923	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	┼
	0,23		-	ļ		+
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	╁
854	1924					T
0.5.5	5020	01,02 (Colon, High Met vs. Colon, Low Met)	8	21		2
855	5838					-
		01,02 (Colon, High Met vs. Colon, Low Met)	1	9		-
856	2062	( a see ), sugar see ( a colon, see inter)	1	9		9
						$\vdash$
İ		13,14 (bFGF Treated HMVEC vs. VEGF-Treated HMVEC) (bFGF			<u> </u>	
		Treated HMVEC vs. VEGF-Treated HMVEC) 03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		6
		01,02 (Colon, High Met vs. Colon, Low Met)	8	19 39		2
857	447388	- (		39		<del>  '</del>
			1	一十		一
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	142	4	38.1	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	18	0	20.57	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	18	0	15.39	
858	12419	15,16 (Normal Colon vs. Colon Tumor Tissue)	142	2	75.05	
000	12419		+ -			<u> </u>
_		03,04 (Breast, High Met vs. Breast, Non-Met)	0	15		1.5
859	3224	, John Steeling Avenue	+ -	13		15
			<del>                                     </del>	$\dashv$		
		01,02 (Colon, High Met vs. Colon, Low Met)	17	2	7.84	_

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	
860	5474					Τ
						T
		03,04 (Breast, High Met vs. Breast, Non-Met)	5	17		
		01,02 (Colon, High Met vs. Colon, Low Met)	1	8		П
861	3522					
		01,02 (Colon, High Met vs. Colon, Low Met)	3	12		+
862	731785					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)			6.00	_
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.09	╀
863	3765	15,10 (Normal Colon Vs. Colon Turnor Hissue)	0	6	-	
		0102/01 W. W. W.				
864	640323	01,02 (Colon, High Met vs. Colon, Low Met)	19	6	2.92	╀
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
865	379105			<u> </u>		_
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		-
866	448029					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	-
867	650476				10.10	
	·	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7		7.51	_
868	640525	15,17 (Normal Colon Tissue vs. Colon Metastasts)	7	0	7.51	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
869	390124	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	_
						-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	18		2
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	5.98	
870	464029	18,19 (Normal Colon Tissue vs. Colon Tumor)	0	8		
070	404029					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9	-	8
871	468109					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
872	21669					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.
873	651088					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)			6.44	
874	2737	10,11 (TOTHIAI COJOR TISSUE VS. COJOR METASTASIS)	6	0	6.44	
		1617(O-1TT)				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	3 4	14		4
875	556421	(		14		3.

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	1 20,
876	452245	13,10 (140111111 COIOII 43. COIOII TUITIOI TISSUC)	12	1	12.08	┼
870	432243			╁─		┢
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	12		11
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	12		11
877	447539	15,17 (Normal Colon 1188de vs. Colon Metastasis)		12		11
	117335			<del> </del>		<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	<del> </del>
		15,16 (Normal Colon vs. Colon Turnor Tissue)	8	1	8.46	-
878	546642	( 13000)		$\vdash$	0.70	╁
				<del>                                     </del>		<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	<b>†</b>
879	236368		·			<b> </b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	271	16	17.9	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	271	0	290.88	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	16	0	16.25	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	9	1	10.29	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	9	0	7.69	
880	644523					
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	6	0	6.09	
881	729173					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7
882	8315					
	-					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
883	450463					
		151( OL1C 1		- 10		
884	650856	15,16 (Normal Colon vs. Colon Tumor Tissue)	31	13	2.52	
004	030830					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.00	
885	648109	10,17 (Colon Tuniol Tissue vs. Colon Metastasis)	0	0	6 09	
000	010103					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
886	726644	. ,		Ť	5.57	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	18	0	18.28	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	18		17.
887	727224					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.5
888	557906					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	
889	502683					
				1		

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	12		11.35
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	12	0	12.19	11.33
891	647952	10,17 (Colon Tunior Tissue 15: Colon Frenchista)	12	Ť	12.17	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
892	639991					
902	725246	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
893	735346					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0,05	5.68
894	102655					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	0	14.8	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	0	15.03	
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	33		8.46
895	553629					<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	17		4.19
		15,17 (Color Turnor Fissue vs. Color Metastasis)	4	17		3.96
896	1609	12517 (1707) Mar Colon 215000 15. Colon 175000 Marsh		17		2,50
		01,02 (Colon, High Met vs. Colon, Low Met)	3	58		20.96
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	2	4 88	
-		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	3	5.37	
	644004	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	14	3	4.74	
897	641884					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
898	648872					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
899	644242					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
900	63559			_	2.2	
	!					
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	12		6.15
901	550108					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	37		5.75
002	274206	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	19		3
902	374306					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32
903	5838					J.D.
		01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/
904	645530					
			1			ĺ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
905	649732					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
906	649143					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
907	7571					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.0
908	4572					
700						
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	11		5.0
909	2147	1-2-5-7	<del>-   -</del>			
707	2171		<del>                                     </del>			<b></b>
		01,02 (Colon, High Met vs. Colon, Low Met)	31	6	4.77	<b></b>
		03,04 (Breast, High Met vs. Breast, Non-Met)	12	2	5.85	
910	462659	05,04 (Diedst, High Met Vs. Diedst, Non Mety	1.2		2.03	
710	402037					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
911	727723	15,10 (140/1112/1 00/01/143. 00/01/14/10/11/13320)	<del>-   '</del>	l	7.1	
	12/123					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.05	5.0
912	2636	15,10 (110/11141 - 65/01/15) - 65/01/14/10/11/15546)		Ť		
712	2030		···			
		08,09 (Lung, High Met vs. Lung, Low Met)	7	1	9.78	
		01,02 (Colon, High Met vs. Colon, Low Met)	8	19	70	2.:
913	500959	101,02 (Colon, Thigh Met vo. Colon, Low Met)		17		
	500252		<u> </u>			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	$\vdash$
914	3428					
	3.20					
-		03,04 (Breast, High Met vs. Breast, Non-Met)	20	0	19.51	
915	734929					
				$\vdash$		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.0
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
916	453592	10,1. (Colon Tainor Tiboue 15. Colon Michaelasia)		۱	5.07	<del> </del>
710	700016					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	6	3.35	<del>                                     </del>
917	15414	15,15 (Atomiai Colon 15, Colon Tumor Hosac)		<del> </del>	2.33	
217	13414			$\vdash$		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.3
918	648959	10,17 (Normal Colon Tissue vs. Colon Metastasis)	-   1	7		-0
710	040939			$\vdash\vdash$		<b></b> -
		15 17 (Normal Colon Tissue va Colon Mataria)	7	_	7 51	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	/	0	7.51	L

## The first field that the first state of the state of the state state with the state of the state

## Table 5

Table 5			1 .			T = -
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
919	453470	Language Control of the Control of t				
						ļ
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	1	12.88	
220	640272	15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	
920	649272					
		16 17 (Color Turner Tierre va Color Meteotoxic)	8	0	8.12	
921	1699	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	°		0.12	
921	1099					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10		9.46
	ļ	03,04 (Breast, High Met vs. Breast, Non-Met)	37	12	3.01	7.70
922	649719	03,04 (Bleast, High Wet vs. Bleast, Non-Wet)		12	3.01	<b></b>
722	043713					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
923	562805					
			V.V. 114			
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
924	452204					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	3	13		4.29
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
925	549178					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	10		9.85
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32
926	639177					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
927	562550					
						7.01
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
928	561807					
	<u> </u>	16.12 (Color Transactions Color Metados)		8		7.00
929	641373	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	1	l °		7.88
929	041373					
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	0	16.1	
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	15	3	5.28	
930	514418	15310 (Normal Colon VS. Colon Tuniol Tissue)		Ť	5.20	
	<b>-</b>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8 86
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.38
931	567078		,			
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	1	11.17	
932	643061					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
933	549160					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
934	449269					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	26	7	3.99	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	26	5	5.5	
935	453082					<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	ļ
-026	110105	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<del>                                      </del>
936	418135					
		16 17 (Color Trans Tierra va Color Meteotoria)	6	23		3.77
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	23		21.43
937	2783	15,17 (Normal Colon Tissue vs. Colon Metastasis)		23		21.7.
937	2703		<del></del>	$\vdash$	·	<del> </del>
		01,02 (Colon, High Met vs. Colon, Low Met)	27	6	4.15	
		03,04 (Breast, High Met vs. Breast, Non-Met)	5	40		8.2
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
938	549435					
						<b> </b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	2	7.4	
939	446614					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
940	449477					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
941	454380					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	14		4.42
942	450914					
				<u> </u>		5.60
0.10	72.000	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
943	736860	<u> </u>		-		<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	·	5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1 6	0	6.09	3.00
944	727224	10317 (Coloii Tulloi 1138de vs. Coloii Medastasis)	<del></del>	<u> </u>	0.07	
	12/22-					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
945	644242					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
946	562550					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
947	649148					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		•		. I		
948	375889			<del></del>		
948	375889				-	
948	375889 449437	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs Colon Tumor Tissue)	14	3	4.93	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	12		3.94
950	449044					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
951	555318					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	12		5.91
952	456764	10,17 (00011 101101 110000 10. 001011 11011011010)		1		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	
052	11567	15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	1	15.03	ļ
953	11567				ļ	
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
954	3522					
		01,02 (Colon, High Met vs. Colon, Low Met)	3	12	<u> </u>	4.34
955	456528	01,02 (Colon, High Met vs. Colon, Low Met)		12	<u> </u>	4.34
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
956	639142					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
957	446371	15,17 (Normal Colon 1155ac vs. Colon Metastasis)		Ť	0.57	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	ļ
958	554742	<del> </del>		<b>-</b>		<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
959	448029					
2.42		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	
960	551380					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14		3.45
961	551527					
062	720205	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
962	729295	<del> </del>				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
963	349744					
		16.17 (Color Tomor Times Color Vision				F 0:
964	648996	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
771	V.0770			$\vdash$		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
965	447126			$\vdash$		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	5	4.02	
		<del></del>		للنب		

## Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	5	20		3.94
966	730866					
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	8	0	8.12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
967	420686					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
968	451753					
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	1	12		11.88
		15,16 (Normal Colon vs. Colon Tumor Tissue)	4	15		3.55
969	451380					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	20	7	3.02	
970	645530					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
971	554703					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9		8.38
972	562835					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9		8.38
973	732764					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
974	556216					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
975	728779					L
		<u> </u>				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6 09	
976	414739					<u> </u>
]	i 	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	14		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	14		6.52
977	551514					
977	551514					
977	551514	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	13		12.8
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	1 8	13	8.46	12.8
977	551514 550107	<del>                                     </del>	<del></del>		8.46	12.8
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1		12.8
978	550107	<del>                                     </del>	<del></del>		2.72	12.8
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1		12.8
978	550107	15,16 (Normal Colon vs. Colon Tumor Tissue)  15,16 (Normal Colon vs. Colon Tumor Tissue)	36	1 14		
978	550107	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1		7.57

Table 5

Table 5			<del></del>			
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
	<del> </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	}
981	562550	13,17 (Normal Colon Tissue vs. Colon Niciastasis)		1	0.44	
	1 302000	<del> </del>		†	<b></b>	<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
982	549722					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
983	640525			ļ		
	<u> </u>		<u>_</u> _	<b> </b>		<b> </b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<b> </b>
984	455542	15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	<del> </del>
204	433342					<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	6		5.68
985	9436					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	15		3.69
986	380284					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9.51	
007	55(2(0	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	1	9.66	
987	556260					<del></del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
988	650476	1947 (Colon Famor Fissac 13. Colon Pretastasis)		Ť		3.51
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
989	554500					
ē						
·····	ļ	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
990	422375					
	<del> </del>	15 17 Olympia Color Time of Color Manager		<del> </del> _		5 12
991	456528	15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
771	430328					
	<b></b> -	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
992	644190					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
993	554080					
004	546205	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
994	546705		<del></del>			
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
995	558337	10,17 (Colon Tunor Tissue vs. Colon Wetustasis)		<del>                                     </del>		3,91
	-3025/					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
996	449269					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	26	7	3.99	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	26	5	5.5	
997	645799					

able 5	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
998	456506					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	17	7	2.61	
999	219416	15,17 (Normal Colon Tissue vs. Colon Metastasis)	- 1'		2.01	
999	218416					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1000	455820					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1001	554703					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0_	9	<u> </u>	8.38
1002	650204					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	
1003	456808	15,17 (Normal Coloir Tissue vs. Coloir Wetastasis)		Ť		
1003	430000					<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	42		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	42	1	42.65	
1004	420686					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
1005	378373			<del>                                     </del>		-
		Color (Color Metasteria)	0	7		6.89
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1006	463824	15,17 (Normal Colon Tissue vs. Colon Metastasis)	-	<u> </u>		
1000	403624					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1007	24939			<u> </u>		ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.5
1008	556561			-		<u> </u>
		1/ 17/G 1 T Tiene un Celen Metesteus)	1	10		9.8
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	10		9.3
1009	380406	13,17 (Normal Colon Tissue vs. Colon Metastasis)				
1009	300400					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	12		3.9
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	3	5.64	
1010	456764					
				<u> </u>		ـــــــــــــــــــــــــــــــــــــ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	1	15.03	
	ļ	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	1 -
1011	725703			├		+
	<b>_</b>	15.16 (Namuel Colon or Colon Turner Tirene)	0	9		8.5
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	1 3.2
1012	185465	10,17 (Coloil Turnor Hissue vs. Coloil Miciastasis)		Ť		1
1012	100700			T	t	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	14		7.17
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14		3.45
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	25	9	2.81	
1013	5830					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1014	539955					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	24	0	24.37	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	3	47		15.51
1015	640747					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1016	500630			-		
		Time Classic		0	6.09	<u> </u>
1015	440511	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	0.09	<del> </del>
1017	448511					<del>                                     </del>
		16.17 (Color Tyreer Ticque va Color Metectoris)	4	26		6.4
	<u>.                                    </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	26		8.07
1010	405073	13,17 (Normal Coloil Tissue vs. Coloil Metastasis)		20		0.07
1018	403073			-		<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1019	641439	15,75 (Nothing Color to Color				
1015	047.55					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1020	406092					
						<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	3	4.29	
1021	559806			<u>L</u>		
						1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1022	380284			<u> </u>		-
				<u> </u>	0.51	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9.51	<del> </del>
	560700	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9_	1	9.66	+
1023	560700			<del>                                     </del>	<del>                                     </del>	+
		16 17 (Color Turner Tienne un Color Matentagis)	0	6		5.91
1024	552879	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		1		3.91
1024	332819			<del>                                     </del>	<del>                                     </del>	<del>                                     </del>
		16.17 (Colon Tumor Tissue vs. Colon Metastasis)	2	14		6.89
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	1
1025	640590	15,15 (Ivorniai Colon vs. Colon Tunior Hissac)		Ť	1	<b>†</b>
1023	040590			<b>†</b>		<b>†</b>
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
	<u> </u>	15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
1026	641683					
	†	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	

Table 5

Table 5						T
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1027	648934					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1028	557948					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
1029	377094					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
1030	449617					
1050	447017					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	18		2.53
1021	079	10,17 (Coloii Tunioi Tissue vs. Coloii Metastasis)				
1031	978			$\vdash$		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	67	20	3.6	
		The same of the sa	67	28	2.53	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	78	23	3.31	
		03,04 (Breast, High Met vs. Breast, Non-Met)	/8	23	3.31	
1032	607430					
					6.24	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1033	641837					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	16	0	17.17	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	31	0	31.48	
1034	449750			L		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	28		27.57
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	28		6.52
1035	646780					
						ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	ļ
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1036	546642					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
1037	642906					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1038	552879					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	14		6.89
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	
1039	644205					
	T	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1040	506744					
	<b>1</b>					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
i -	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1041	557797					
<b>─</b> ─	1			1		1

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ.	0202	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	16		5.25
1042	640356	10,17 (Colon Tunios Tibers				
1042	040330					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
1042	462650	15,17 (Nothiai Colon Tissue vs. Colon Metastasio)				:
1043	462659					
		C. L. Turner Tienne)	7	0	7.4	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.51	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		<del>-                                    </del>	7.51	
1044	645633					
					2.51	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1045	237288					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1046	454343					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.16	
1047	386543					
1017	3300 10					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	23		7.55
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	23		7.14
1048	446404	13,17 (Normal Colon Hisbac to Celon Fitter)				
1048	440404			<u> </u>		
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	23	0	19.66	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	73	0	78.36	
	<u> </u>		23	0	26.28	<del>                                     </del>
· _		18,19 (Normal Colon Tissue vs. Colon Tumor)	73	1	77.16	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	<del>                                     </del>	17.10	<del></del>
1049	456528			-	<u> </u>	<del></del>
	ļ			7		6.62
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	<del>  '</del>	ļ	0.02
1050	456528			├		<u> </u>
_				<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1051	452781					
				ļ		-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
1052	551671			<u> </u>		ļ
				<u> </u>		
<del></del>		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6 52
1053	644242					
	<del>                                     </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<u> </u>
	1	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1054	561892					
1004	301072					
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.5
1055	450420	13517 (HOTHER COION HISSEC VS. COION HICESEED)		1	<del>                                     </del>	1
1055	450429			+-	†	1
	-	16.12 (O. L. T Tirana v. Callar Matastagia)	10	1	10.16	+
	+	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	1	13.95	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	- 13	+ 1	13.73	1-
1056	533588		I	l	1	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1057	553877		<del></del>			
					<u> </u>	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
1058	650195					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1059	193486					
-						
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1060	650195					
				1		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1061	562835	, , ,				
					<b> </b>	<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9	<b> </b>	8.38
1062	736816	, , , , , , , , , , , , , , , , , , , ,		<u> </u>		
					<del>                                     </del>	
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	9	0	9.14	
	<del></del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
1063	403632			·		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1 2	11		5.12
1064	390124					
						<del> </del>
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	8		7
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	18		2.79
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	5.98	
1065	390124					<b>-</b>
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	8		7
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	5.98	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	18		2.79
1066	422687					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	10		9.46
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.16	
1067	394413					
		18,19 (Normal Colon Tissue vs. Colon Tumor)	21	2	12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	113	3	39.81	
	**	15,17 (Normal Colon Tissue vs. Colon Metastasis)	113	0	121.29	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	21	0	17.95	
1068	549178		<del> </del>			
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	10		9 85
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32
1069	453079					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	
1070	463824					
		<del></del>				
				t		

6.34 6.44	B/A
	<u> </u>
6.44	T 7
	<del> </del>
<del> </del>	
<del> </del>	<del>   </del>
	5.68
6.09	-
<u> </u>	
	8.46
14.8	<del> </del>
	<u> </u>
	6.21
6.44	
	12.11
	6.62
4.02	
	3.94
	9.32
)	9.85
)	4.92
)	9.32
	6.52
	5.91
3.17	
6.44	
4.23	
) [	8.38
,	8.51
3 6 3 6 3	15.03 14.8 14.8 14.8 14.8 16.44 17. 18. 19. 19. 19. 19. 19. 19. 19. 19

Table 5

Table 5						
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1086	451243			<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6 52
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	<u></u>	5.68
1087	452506			<u> </u>		<u> </u>
				<u> </u>		<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	<u> </u>	5.91
1088	554703	<u> </u>				<u> </u>
				<u></u>		<u></u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9		8.38
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
1089	449580					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	4	4.23	
1090	3316					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5 91
1091	97507					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	9		8.51
1092	556216					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1093	185401					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	63		64.57
1094	3758					l
		03,04 (Breast, High Met vs. Breast, Non-Met)	17	5	3.32	
1095	95700					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	21		21.52
1096	2478					
		03,04 (Breast, High Met vs. Breast, Non-Met)	17	5	3.32	
1097	550267					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	4	4.03	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	1	15.85	
1098	185652					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
1099	55798					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
1100	5078		<u> </u>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1101	9784					
		<del></del>				<u> </u>
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
1102	2245		<del>-   -</del>		<del></del> -	<del></del>
		<del></del>				

Table 5

Table 5						
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
				<u>L</u>		
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	L
		01,02 (Colon, High Met vs. Colon, Low Met)	12	27		2.44
1103	11606			<u> </u>		<u></u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
1104	2245			<u> </u>		<u>l</u>
				<u> </u>		
L		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	
		01,02 (Colon, High Met vs. Colon, Low Met)	12	27		2.44
1105	551172					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10		9.46
1106	729175					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1107	6317					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1108	2478					
		03,04 (Breast, High Met vs. Breast, Non-Met)	17	5	3.32	
1109	4727					
		03,04 (Breast, High Met vs. Breast, Non-Met)	19	0	18.54	
1110	185598					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2
1111	736349					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1113	189561					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
		08,09 (Lung, High Met vs. Lung, Low Met)	1	14		10.02
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9		8.38
1114	728131					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1115	560984					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	- <del></del>	5 91
1116	549945		1			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	2	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	2	6.34	
1117	554785					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
1118	554785					
		<del></del>				

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLSI	Library Tan A,D		-	73/B	1 27.73
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8	<b></b>	7.88
1119	551235	10,17 (Colon Tuthor Tissue vs. Colon Metastasis)	<del></del>	l °		7.00
1119	331233			<del> </del>		├──
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		7		6.89
1120	2634	10,17 (Colon Tuntor Tissue vs. Colon Wetastasis)	<del></del>	<del>                                     </del>		0.05
1120	2054			╁┈─		┼
		03,04 (Breast, High Met vs. Breast, Non-Met)	48	0	46.83	<del>                                     </del>
1121	548858	03,04 (Breast, Trigit Wet vs. Breast, Non-Wet)		<del>                                      </del>	40.03	<del>                                     </del>
1121	348838			<del> </del>		<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		11	<b></b>	10.83
1122	15625	10,17 (Coloit Turior Tissue vs. Coloit Wetastasis)	<del>-                                    </del>	1		10.6.
1122	13023			<del></del>		├──
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	17	0	18.25	<del>                                     </del>
		15,16 (Normal Colon Vs. Colon Tumor Tissue)	17	1	17.97	<del>                                     </del>
1123	649259	13,10 (Normal Colon vs. Colon Tunor Hissac)	<del></del>		17.57	├──
1123	049239					
	<del></del> -	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	├──
1124	550267	13,10 (Normal Colon vs. Colon Tunior Tissue)	<del></del>	l <del>"</del>	<del></del>	├
1124	330207			-	<b> </b> -	<del></del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	1	15.85	<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	4	4.03	+
1125	7436	13,17 (NOTHIAI COIOII TISSUE VS. COIOII METASTASIS)	- 13	-	7.03	├
1123	7430			<del>                                     </del>	<u> </u>	<del> </del>
	L	03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	├──
1126	451794	05,04 (Bleast, Fight Met 13. Bleast, Foll-Met)	<del></del>	<u> </u>	0.70	
1120	431174	<del> </del>				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	0	15.85	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	1	16.1	<del>                                     </del>
1.127	5744	10,17 (1407) and Colon Fisher 10. Colon Fisher Colon	<del></del>	H	1311	<del> </del>
1.127	3744					$\vdash$
		01,02 (Colon, High Met vs. Colon, Low Met)		6		6.5
1128	3516	01,02 (Colon, Tright Net vs. Colon, Low Net)	+ <u>*</u>	Ť		
1120	3310			_		
		01,02 (Colon, High Met vs. Colon, Low Met)	5	17		3.69
1129	730555	(Coon, 1.1gn 1.cc to. Colon, 20 v 1.cc)				-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1130	3085					
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	11		5.64
1131	638854		<del></del>			
						<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	32	11	3.07	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	32	0	34.35	
	<del></del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
1132	7379		<del></del>			
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1133	185562	, , , , , , , , , , , , , , , , , , , ,	- + -			<del></del>

SEQ	CLST	Library Pair A,B	A	В	<u>A/B</u> _	B/A
<u> </u>		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
1134	452491	00,0 . (2.0.0.), - 2				
1154	432471					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	
1105	(46248	13,10 (Notifial Colon vs. Colon Tunior Fisade)				
1135	646248					
		Times of Color Metasteria	6	0	6.09	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	·	<u> </u>		
1136	6056					
				<del>_</del>	6.92	
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
1137	643103					ļ
						<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	ļ
1138	6923					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
1139	6923					Ī
1137	0,25					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
1140	001	05,04 (Bleast, Tright Mee 18. Bleast, 11611 515)				
1140	901					
		102 04 (D IVl. Mature Dreast Non Mat)	56	5	10.93	
		03,04 (Breast, High Met vs. Breast, Non-Met)		ا ا	10.55	<u> </u>
1141	901			-		
				5	10.93	<del> </del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	56	3	10.93	<del>                                      </del>
1142	367					<del>                                     </del>
						226
		08,09 (Lung, High Met vs. Lung, Low Met)	30	99	2.44	2.36
_		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	4	3.44	
		03,04 (Breast, High Met vs. Breast, Non-Met)	105	24	4.27	1
		01,02 (Colon, High Met vs. Colon, Low Met)	19	97		5.53
1143	4043					
		01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76
1144	3299					
	<del>                                     </del>	08,09 (Lung, High Met vs. Lung, Low Met)	10	1	13.97	
	<del>                                     </del>	03,04 (Breast, High Met vs. Breast, Non-Met)	20	3	6.5	
1145	11881	00,00 (60,000,000,000,000,000,000,000,000,000,				
1143	11001			1		
	<del> </del>	03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5 85	
1111	0112	03,04 (Bleast, flight Met Vs. Bleast, Mon-Met)	<del>                                    </del>	Ť		+
1146	9113			<u> </u>	<del>                                     </del>	+
	ļ	The second state of the se	0	7	$\vdash$	7.59
		01,02 (Colon, High Met vs. Colon, Low Met)		+ -	<del>                                     </del>	1.35
1147	185460			+	<del> </del>	+
	<u> </u>			1	-	10
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	16		16.4
1148	185716			1		
						<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	6	<u> </u>	6.15
1149	5753				<u> </u>	
	<del>                                     </del>					
	_ <del> </del>	01,02 (Colon, High Met vs. Colon, Low Met)	0	10	I	10.8

Table 5

Table 5	CLST	Library Pair A,B	A	В	A/B	B/A
<del></del>		Littary I all A,D	A	D	AD	D/A
1150	24939				<del> </del>	<b>├</b> ──
				L	<u> </u>	ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8	ļ	7.57
1151	649684			<u> </u>	<u></u>	<u> </u>
						<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
1152	642109					1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1153	15035	<del>                                     </del>				1
	10 033	<del> </del>				<del></del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	<del> </del> -
	<u> </u>					<del> </del>
	6405.74	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8 59	├
1154	649354					<del> </del>
				ļ		├
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	L
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<u>L</u>
1155	4465					
		01,02 (Colon, High Met vs. Colon, Low Met)	4	14		3.79
1156	647952					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1157	455601	te, re (remain colon to colon rame)	<del>-</del> -	Ť	0.5.	
1157	433001			<del>  -</del>		<del> </del>
		16 17 (Colon Turnor Tiguro ya Colon Matartaria)			8.12	<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	<del></del>	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
1158	641901					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<u> </u>
1159	446878					
						<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	12		11.82
1160	7436					Ī — — —
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
1161	2245					
		01,02 (Colon, High Met vs. Colon, Low Met)	12	27		2.44
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	
1162	3531	05,04 (Bleast, High Met Vs. Bleast, Non-Met)	13		12.00	
1102	5551					
		01,02 (Colon, High Met vs. Colon, Low Met)	<del></del>	17		6.14
	0605	101,02 (Colon, High Met vs. Colon, Low Met)	3	17		6.14
1163	9625	<del> </del>				
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	12		12.3
1164	727489			]	·	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
1165	159925					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	22		22.55
		· · · · · · · · · · · · · · · · · · ·				

Table 5

Table 5			<del></del>	Τ=	<del></del>	T = '
SEQ	CLST	Library Pair A,B	A	B	A/B	B/A
1166	645210			<del> </del>	<u> </u>	
	<u> </u>			<b>├</b>	10.55	ļ
	ļ. <del></del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	10	1	10.57	<u> </u>
11.5	1.77(00	15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	<b></b>
1167	157629	<del> </del>		<del> </del>	ļ	
		02.04 (P		10		(15
1160	0275	03,04 (Breast, High Met vs. Breast, Non-Met)	3	18		6.15
1168	8375	<del></del>				
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	<del> </del>
1169	4319	05,04 (Bleast, High Met Vs. Bleast, Noh-Met)	<del></del>	<del>                                     </del>	0.63	
1109	4319		<del></del>			<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	14	0	13.66	<del> </del> -
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6	13.00	6.5
1170	4045	01,02 (Colon, riigh wet vs. Colon, Low Met)	_ <del></del> -	<del>                                     </del>		0.5
1170	4043	<del> </del>	<del></del> _	<del> </del>		<del> </del>
		01,02 (Colon, High Met vs. Colon, Low Met)	2	11		5.96
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	3.50
1171	185642	05,07 (Bloust, Fight Met 13. Bloust, 1107 Met)		<del>                                     </del>	7170	
	103012	<del> </del>		1-		
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
1172	7436	one (States, Ing.) The total states		<u> </u>		
11.12	7.50	<del> </del>		<del>                                     </del>		
		03,04 (Breast, High Met vs Breast, Non-Met)	9	1	8.78	
1173	3531					
			<del></del>			
		01,02 (Colon, High Met vs. Colon, Low Met)	3	17		6.14
1174	644776					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
1175	8354					
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
1176	2099					
		03,04 (Breast, High Met vs. Breast, Non-Met)	27	7	3.76	
1177	449956					
	i	<u> </u>				
	l	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	
1178	649106	<u> </u>		<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1179	452414					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	17	0	17.97	<u> </u>
1100	700710	15,17 (Normal Colon Tissue vs. Colon Metastasis)	17	0	18.25	
1180	732712	<del></del>				
		16.17.60.1 Tomas Times of Col. Miles (1)			7.11	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	6.62
1191	195562	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1181	185562	<del></del>	<del></del>	<del>  </del>		
		03,04 (Breast, High Met vs. Breast, Non-Met)	- 0	9		9.22
1182	3516	05,07 (Dicast, High Mct vs. Dicast, MON-Met)	<del></del>	"		7.64
1104	2210	1	L	L _ [	l '_	l

Table 5	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLSI	Library Pair A,D		B	A/B	D/A
	<u> </u>	<del> </del>		ļ		<del> </del>
	<u> </u>	01,02 (Colon, High Met vs. Colon, Low Met)	5	17		3.69
1183	185562			ļ		ļ
				<u> </u>		<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
1184	185460	<u> </u>				<u> </u>
						<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	16	]	16.4
1185	10947					Ţ
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	0	7.81	
1186	452856					<u> </u>
	<del>                                     </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	1
1187	558767			<del>                                     </del>		<del>                                     </del>
1107	330707	<del> </del>				<del>                                     </del>
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
1188	15035	13,10 (Normal Colon vs. Colon Tumor Tissue)	<del>+-</del> -	┼╌	0.40	<del>                                     </del>
1100	13033	<u> </u>				├
		15 17 Ol and Color Times on Color Materials		<del>  , </del>	9.50	<b>├</b> ──
	ļ	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)		0	8.46	<b>├</b>
1189	556421			<u> </u>		<del> </del>
	<u> </u>			<u> </u>		<b></b> _
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	<b> </b>
1190	7082					<b>!</b>
					Ĺ	Ļ
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	0	9.76	<u> </u>
1191	452523	<u> </u>				
						Ĺ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
1192	3242					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	17		9.22
1193	6660					
		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	,
1194	547					
		01,02 (Colon, High Met vs. Colon, Low Met)	35	67		2.08
	[	03,04 (Breast, High Met vs. Breast, Non-Met)	90	30	2.93	1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10		9.46
1195	121213					
	121212	<del></del>		<del>                                     </del>		<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	<del> </del>
1196	4378	10,17 (Colon Tulliol 1188tic vs. Colon Mictastasis)	<del>+</del> -	l -	0.12	
1190	4378	<del></del>	<del></del> -	-		<del> </del>
	ļ	16 17 (Color Trans Trans Color Maria	<del>+</del> -			5 10
<del></del> -	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	11		5.42
	ļ	03,04 (Breast, High Met vs. Breast, Non-Met)	1	8		8.2
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	
1197	185554	<u></u>				
	ļ	<u></u>		$\sqcup$		
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
1198	185482			لــــا		
	]		1	1 1		ł

Table 5

Table 5	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	13		13.32
1200	66017					1
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2
1201	403111					1
				_		<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<u> </u>
1202	3224					
		<del> </del>				
		01,02 (Colon, High Met vs. Colon, Low Met)	17	2	7.84	
1203	966					
٠						
		01,02 (Colon, High Met vs. Colon, Low Met)	22	47		2.32
		03,04 (Breast, High Met vs. Breast, Non-Met)	50	19	2.57	
1204	3639					
		03,04 (Breast, High Met vs. Breast, Non-Met)	12	3	3.9	
1205	5388					
		03,04 (Breast, High Met vs. Breast, Non-Met)	5	21		4.3
1206	3299					
	<del></del>					<u> </u>
		08,09 (Lung, High Met vs. Lung, Low Met)	10	1	13.97	
		03,04 (Breast, High Met vs. Breast, Non-Met)	20	3	6.5	<u> </u>
1207	23760					<u> </u>
				-		<u> </u>
1200	720204	03,04 (Breast, High Met vs. Breast, Non-Met)	- 4	34		8.71
1208	729384					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	$-\frac{1}{7}$	0	7.11	<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	- 10	7	7.11	6.62
1209	46559	15,10 (Normal Colon vs. Colon Tullor Tissac)	<u>_</u>			0.02
1207	40337					<del>                                     </del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	30		15.37
	<del></del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	13.57
1210	449750					<b></b> -
				$\vdash$		t
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	28		6.52
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	28		27.57
1211	735936					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
1212	607430					
T						
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1213	452856					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1214	557903					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9	-	8.51
1215	453112	<u> </u>				L

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	13		4.27
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	13		4.04
1216	645900					
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	7	0	7.07	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
1217	415114			<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	0	10.57	
1218	418763					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
1219	2245				<u> </u>	
		02.04 (D		<u> </u>	10.60	<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	
1220	403668	01,02 (Colon, High Met vs. Colon, Low Met)	12	27		2.44
1220	403008					
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	3	4.4	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	13	4.4	6.15
1221	15427	15,10 (Normal Colon Vs. Colon Tunor Tissue)		13		0.13
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
1222	555714			<u> </u>		1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	23	11	2.21	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	23	7	3.53	
1223	555830					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	16	2	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	2	8.46	
1224	4620					
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
1225	171511					
		1/ 17 (C.1 T Time - C.1 - M )			G 11	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	( (2
1226	451401	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1220	731401					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	1	9 66	
1227	447501		7	1	7 00	
,	,501					
<u> </u>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	3	5.37	
1228	460445	( Colon Manuala)	13		3.37	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1229	375814				-	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	

Table 5

Table :			<del></del>	T		<del></del>
SEQ	CLST	Library Pair A,B	A	B	A/B	B/A
1230	449356			<u> </u>		ļ
<u> </u>				-		
<u> </u>	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	21		2.79
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	21		5.17
1231	468736			<b>-</b>		
<b>}</b>				ļ.,		
	5100.50	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	<u> </u>	5.91
1232	548858	<del></del>		<del>  </del>		<b>}</b> -
		1/17/01 7 7 7		<del> </del>		10.00
1222	2602	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
1233	3693			-		<del> </del>
		02 04 (Proced High Maters Proced New Mat)		-		(15
1224	(42072	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
1234	642973	<u> </u>		<del> </del>		<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7 1 1	<del> </del> -
1235	561180	10,17 (Coloit Tullor Hissue vs. Coloit Metastasis)	<del></del>	<u>'</u>	7.11	
1233	301180	<del> </del>				<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	<del> </del>
1236	453708	13,10 (Normal Colon vs. Colon Tumor Tissue)			3.81	
1230	433708			-		<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	19		18.71
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	19		17.7
1237	645305	13,17 (Normal Colon Tissac vs. Colon Metastasis)	<del></del>	12		17.7
1207	0.0000	<del> </del>				<del> </del> -
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<del> </del>
1238	463487	10,17 (Conta) Colon 110500 Vol. Colon (Voluments)		Ů	7.51	
1250	102107	<del> </del>		$\vdash$		<del>                                     </del>
	<u></u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
1239	11131		<del>-   -</del>			
						<del> </del>
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1240	561807					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
1241	452800					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	13		4.04
1242	372960					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	18		2.79
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	33		5.2
1243	449317					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	5	20		3.73
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	20		2.46
1244	730759					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1245	9113					
124	6202-0	01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59
1246	630259	<u> </u>				

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1247	3516					
		101 02 (Colon High Mat vs. Colon Low Mat)	5	17		3.69
1248	447494	01,02 (Colon, High Met vs. Colon, Low Met)	<del>-   '</del>	1/		3.09
1240	44)494			-		<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	26	8	3.44	
1249	554500					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1250	639662					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1251	421	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1231	421					
	<del> </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1252	736014					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1253	643061					
						ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1254	9113	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
1234	9113			$\vdash$		
·	<del></del>	01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59
1255	650856					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1256	476223					
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	2	19	7.51	9.4
1257	737088	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1437	131066					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1258	449512					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	14		3.26
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	14		13.79
1259	449457					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)				( 52
1260	521901	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1200	321901					·
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1261	175799					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	3	4.93	
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	1	9		8.91

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1262	550108					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	19		3
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	37		5.75
1263	203605		<del></del>	+	<del>                                     </del>	3.7.5
		<del> </del>		<del> </del>	<del> </del>	<del> </del>
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	17	7	2.57	<del> </del>
1264	450429	13,10 (Normal Colon Vs. Colon Tulior Tissac)	<del></del>	<del>                                     </del>	2.57	<del> </del> -
1204	430429			1-	<del> </del>	<del> </del>
	<del> </del>	15 17 (Named Color Tierra or Color Materia)	<del></del>	├	12.05	<del> </del> -
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	1	13.95	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	<del> </del>
1265	2478			<b>!</b>		<b> </b>
				<u> </u>	<u> </u>	
		03,04 (Breast, High Met vs. Breast, Non-Met)	17	5	3.32	<b> </b>
1266	644099					
				L	L	<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1267	552614					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1268	452523					
				-		
	<del></del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
1269	446789	1.000	<del>-   :</del>		- 5.5.	<del></del>
		<del> </del>	<del></del>			<del> </del>
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	16	5	3.38	
1270	515631	15,10 (Normal Colon Vs. Colon Tumor Tissue)		-	3.56	
1270	313631	<del></del>				<del> </del>
<del>, , , ,</del>		15,16 (Normal Colon vs. Colon Turnor Tissue)	0	6		5.60
1271	452522	13,10 (Normal Colon vs. Colon Tumor Tissue)		-	<u> </u>	5.68
1271	452523	<del></del>		<del> </del>		
	ļ	10.00		<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
1272	640116			<b> </b>		
	ļ					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	19	0	20.39	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	4	5.02	
1273	9113					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59
1274	562221					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
1275	455972					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1276	449137					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	18		5.68
1277	5078	Colon to Colon Latitud Library	<del> -</del> -			2.00
1211	20/0					
	<del></del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0			5.60
1270	5079	1.5,10 (Normal Colon vs. Colon Tumor Hssue)		6		5 68
1278	5078	<del> </del>				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68

Table 5

CLST	Library Pair A,B		l D	LAZE	
	Library Fall A,D	A	B	A/B	B/A
4016			<u> </u>	<b> </b>	L
			<u> </u>		<u> </u>
<del> </del>		<del>}</del> -	6	<b> </b>	6.15
402111	01,02 (Colon, High Met vs. Colon, Low Met)	5	14		3.04
403111				<b> </b>	<b></b> -
	15 16 (Normal Colon vs. Colon Turner Tiesua)		-	6.24	<del> </del>
			<del> </del>	<del>}</del>	<del> </del>
562292	15,11 (Totalia Colon Fisher 13. Colon Metastasio)	<del>-</del>	1 "	0.44	<del> </del> -
	<del> </del>		<del> </del>	<del> </del>	<del> </del>
	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	<del> </del>	5.91
403111					
	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
403111					
			L		
	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<u> </u>
	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<b></b>
500959			<b> </b>		<b> </b>
	16 17 Observal Color Times of Color Materials		-	0.50	<b> </b>
				<del></del>	<del> </del>
763	13,16 (Normal Colon vs. Colon Tumor Tissue)		-	8.46	<del> </del>
703			<del> </del>		
	01 02 (Colon, High Met vs. Colon, Low Met)	29	77		2.88
				4.1	2.00
763	(2-tably tage and take a state)	<del>+</del> -			
	03,04 (Breast, High Met vs. Breast, Non-Met)	42	10	4.1	
	01,02 (Colon, High Met vs. Colon, Low Met)	29	77		2 88
500959					
		8	0	8.59	
	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	ļ
452071					
	15 17 Ot 1 C-1 - T' C 1 - M ' )		لـــِـا	115	
	<del> </del>				<u> </u>
468672			<u> </u>		
	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
455492					
	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
639667					
<del></del>					
	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
	l	1 1			
549829	<del> </del>				
	403111 500959 763 763 500959 452071 468672	15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis) 562292  16,17 (Colon Tumor Tissue vs. Colon Metastasis) 403111  15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue) 403111  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis) 500959  15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Vs. Colon Tumor Tissue) 763  01,02 (Colon, High Met vs. Colon, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met) 763  03,04 (Breast, High Met vs. Breast, Non-Met) 500959  15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis)	01,02 (Colon, High Met vs. Colon, Low Met)   5	01,02 (Colon, High Met vs. Colon, Low Met)   5   14	01,02 (Colon, High Met vs. Colon, Low Met)   5   14

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1293	553158			+~	122	Ditt
					<del> </del>	<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
1294	561485					
	-					
	<b>_</b>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1227		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1295	639352			<u> </u>		
	<del> </del>	15 17 0/1 G 1 - T'			<u> </u>	<u> </u>
	<del> </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1296	451401	15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1270	431401			┼		
	f	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9		0.66	
1297	643103	1531 (Total Colon Tissue vs. Colon Metastasis)	9	1	9.66	<del> </del>
				+		<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	<del> </del> -
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1298	468736				1	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1299	218416					
1200	447701	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1300	447501			ļ		
		15 17 (Normal Calar Time Calar Calar				
1301	558371	15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	3	5.37	
	2302/1					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
1302	561794		10	1	5.20	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1303	645065					
1204	451260	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1304	451269					
	<del>_</del> '	15,16 (Normal Colon vs. Colon Tumor Tissue)	12	_		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	3	4.58	
1305	401553	(Commit Colon Hobbe vo. Colon Metastasis)	13	4	3.49	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1306	555276			<u> </u>	5.57	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1307	551617					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9.51	
1200		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1308	463480					
-+		16.17 (Color Town Time Color				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1309	549178			+-	1	15,5
				+	<del> </del>	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10	<del> </del>	9.32
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	10	<del>                                     </del>	
1310	374450	( and the colon from the colon		10	+	9.83
				-	ļ	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		+	<del> </del>	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.51	+
1311	562835	[15,10 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1311	302833					
	<del> </del>	15 17 01 10 1 7				
1212	720555	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9		8.38
1312	730555					
	<del> </del>				1	
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1313	732978					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1314	1609					
						1
		01,02 (Colon, High Met vs. Colon, Low Met)	3	58		20 96
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	2	4.88	1 20 70
0		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	3	5.37	<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	14	3	4.74	<del>                                     </del>
1315	18591			1	1.77	<del>                                     </del>
				<del> </del>		<del>                                     </del>
		08,09 (Lung, High Met vs. Lung, Low Met)	8	0	11.18	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	30	8	3.96	<del> </del>
1316	553158	, ( Tissue)	30	l °	3.90	<del> </del>
						<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.00
1317	470602	(Cook Tamer Tissue vs. Colon Metastasis)		8		7.88
	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	9		8.86
1318	639662	13,10 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	
10.0	057002					
		15 16 (Normal Colon ve. Colon Trans. Time)				
-		15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.4	
1319	644721	13,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1017	074/21					
		15 17 OI				
1220	452202	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1320	453202					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
1321	554655					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1322	641988					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1323	453112					

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
				<u> </u>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	13		4.27
	ļ	15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	13		4.04
1324	550694					
	ļ					
ļ	ļ	15,16 (Normal Colon vs. Colon Tumor Tissue)	21	1	22.2	
1225	640106	15,17 (Normal Colon Tissue vs. Colon Metastasis)	21	7	3.22	
1325	649106					
	<del> </del>	15 17 Olympia Color Tive Color Market		4		
1326	638973	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<u> </u>
1320	036973			-		<del>-</del>
	<del>                                     </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	12	+_	12.10	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	0	12.19	
1327	549911	13,17 (100 mail coloit 1155de 13. Coloit Micastalis)		+ -	7.51	
				<del> </del>	<del>                                     </del>	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1328	648774		Ť	+	╁	3.91
				╁┈╴		<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	10	6.44	+-
1329	549911					<b></b>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1330	639662					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1221	560155	15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1331	560455		_			<u> </u>
		15 17 (Norman) Colon Times of Colon				ļ
1332	735805	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1002	733603		-	├		<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.09	5.68
1333	732712		<del>                                     </del>	l		3.08
			-			
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1334	446663					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	32		2.13
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	32		5.25
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		10.7
1225	226224	18,20 (Normal Colon Tissue vs. Colon Metastasis)	1	8		9.36
1335	226324					
		16.17 (Colon Turner Tisque ya Colon Marcon)				
1336	453016	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1220	100010					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	-		6.00	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.09	5.60
1337	550998	(	0	6		5.68
			1			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
				<u> </u>		2.00

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1338	452414					
	<u> </u>					
	ļ	15,16 (Normal Colon vs. Colon Tumor Tissue)	17	0	17.97	
1000	100-00-	15,17 (Normal Colon Tissue vs Colon Metastasis)	17	0	18.25	
1339	129535			_		
·· <u>-</u> .		15,17 (Normal Colon Tissue vs. Colon Metastasis)	41	1	44.01	<u> </u>
		08,09 (Lung, High Met vs. Lung, Low Met)	2	22	11101	7.87
		15,16 (Normal Colon vs. Colon Tumor Tissue)	41	5	8.67	1
1340	447089					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
1341	447850					7.00
		16.17 (Color Turno Tinna Color Maria				
1342	556216	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
12.42	452522	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1343	452523			<u> </u>		<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	-
1344	44424					
		16.17 (Color Turner Times of La Maria				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	41	90		2.16
_		08,09 (Lung, High Met vs. Lung, Low Met) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	35	40		7.16
1345	648872	125,17 (1401min Colon Fissue vs. Colon Metastasis)	25	90		3.35
4						
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1346	451636	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
25.13						
		16,17 (Colon Tumor Tissue vs. Colon Metastasıs)	1	8		7.88
1347	5078					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		£ 60
1348	403111	20,20 (Novinas Colon Vs. Colon Tullor Tissue)	V	0		5.68
		15 17 01 10 1				
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Turnor Tissue)	6	0	6.44	
1349	648959	15,10 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
					i	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1250		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7 51	
1350	380291			$\dashv$		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1351	380291					
		15 16 (Normal Colon vs. Colon Turner Trans)				
1352	230995	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6 62

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1353	562221	·,-		Ť	T	T-//-
		15,16 (Normal Colon vs Colon Tumor Tissue)	8	0	8.46	
1354	450959			<del>  </del>	0.70	-
				<del> </del>		<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	9		8.51
1355	452833	15,10 (Normal Colon 73. Colon Tumor Tissue)		1		8.31
				<del> </del>		<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1356	550195	Total Colon Florida Vol. Colon Florida Vol.	- i	<del>  '</del> -		0.52
				<del> </del>		<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
1357	448927	13,17 (Normal Colon 1188de vs. Colon Michaelasis)		11		3.12
1337	110727			_		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
1358	551514	13,17 (Normal Colon Hissac vs. Colon Metastasis)		12		3.39
1338	331314					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	12		12.0
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	13	9.46	12.8
1359	549829	13,10 (Normal Colon vs. Colon Turnor Tissue)		1	8.46	<u> </u>
1339	349029			<u> </u>		ļ
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)				5.01
1360	551514	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	0	6		5.91
1300	331314					ļ
		15 16 Oleman Colon va Colon Turner Tierre			0.46	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	10.0
1361	561405	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	13		12.8
1301	561485					
		16.17 (Colon Tumor Tiggue va. Colon Metastague)				6.00
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1362	453846	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1302	433640					
		15 17 (Normal Colon Tienners Colon Metatric)		11		10.05
1363	69863	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	11		10.25
1303	09803					
		OS OO (Lynn a High Makers Lynn Lynn Lynn Mae)		- 22		- 10
		08,09 (Lung, High Met vs. Lung, Low Met)	3	23		5.49
1364	727181	03,04 (Breast, High Met vs. Breast, Non-Met)	3	21		7.17
1304	72/101				_	
		15,16 (Normal Colon vs. Colon Tumor Tissue)				5.60
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	6.00	5.68
1365	454050	16,17 (Colon Tumor Fissue vs. Colon Metastasis)	6	0	6.09	
1303	434030					
		15 17 (Named Calon Tierran Calon Cal		<del></del> -	0.50	
1266	725004	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
1366	725994					
		151601 101 01 7 7				
<del></del> į		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1267	1405	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1367	1495					
		02.04 (D				
		03,04 (Breast, High Met vs. Breast, Non-Met)	31	12	2.52	
1368	5665				2.02	

Table 5

SEQ	CLST		A	B	A/B	<b>B</b> /A
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	0	12.68	
1369	5665					
	<u> </u>					
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	0	12 68	
1370	646146					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		+_	100	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	6.09	<del> </del>
1371	8371	(Normal Colon Model 13. Colon Metastasis)	-   *	0	8.59	+
				1	<b> </b>	<del>                                     </del>
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6	<b>†</b>	6.5
1372	73812					
	<del></del> -	03.04 (Project High Metric Project Nov. Mar)				
1373	4242	03,04 (Breast, High Met vs. Breast, Non-Met)	0	12	<del>                                     </del>	12.3
	12.12				<b></b>	<del> </del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	17	0	16.59	+-
1374	5482		17	<del>                                     </del>	10.39	+
				1		<del>                                     </del>
		01,02 (Colon, High Met vs. Colon, Low Met)	1	8		8.67
1375	5474					
		01.02 (Color High May C. L. Y. M.)		<u> </u>		
		01,02 (Colon, High Met vs. Colon, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met)		8		8.67
1376	5448	03,04 (Bleast, High Met vs. Breast, Non-Met)	5	17		3.48
				-		
		01,02 (Colon, High Met vs. Colon, Low Met)	1	10	<del></del>	10.84
1377	7607			1		10.04
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	3	4.65	
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59
1378	555928	15,16 (Normal Colon vs. Colon Tumor Tissue)	13	4	3.44	
1376	333926					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.60
1379	4046			"		5.68
						-
		01,02 (Colon, High Met vs. Colon, Low Met)	4	14		3.79
1380	554080					
		1/ 17/O 1 T T' C 1				
1381	451092	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
1301	431072					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
1382	551380			-		0.00
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14		3.45
1383	546642					
		16.10.10 to m				
1384	1764	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
1304	1704					
		03,04 (Breast, High Met vs. Breast, Non-Met)	-   27	_	- (50	
		01,02 (Colon, High Met vs. Colon, Low Met)	27	8	6.59 2.88	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B,
1385	650773			1		<u> </u>
	1			+	+	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6 44	+-
1386	644205			╅	1 0 44	-
				+		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		+-	<del>  </del>	+
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.51	
1387	105710	13,10 (Normal Colon Vs. Colon Tumor Tissue)	7	0	7.4	
1367	185718					
	<del> </del>					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.
1388	5538					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	_
1389	7546					<b>†</b> –
						+
		01,02 (Colon, High Met vs. Colon, Low Met)	8	0	7.38	+-
1390	727789			۲Ť	7.58	+
				$\vdash$	<u> </u>	┼
		15,16 (Normal Colon vs. Colon Tumor Tissue)				<del>  _</del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.6
1391	3837	1 syr (coton rumor rissue vs. coton victastasis)	6	0	6.09	<del>                                     </del>
	3037			<u> </u>		
		02 04 (Drough High Max. D.				
1392	380477	03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
1392	380477					
1222		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1393	3299					
		03,04 (Breast, High Met vs. Breast, Non-Met)	20	3	6.5	
		08,09 (Lung, High Met vs. Lung, Low Met)	10	1	13.97	<u> </u>
1394	448853					
						_
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10		9.4
1395	736701					7.7
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	10	0	10.1	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	10.1	
		15,16 (Normal Colon vs. Colon Tumor Tissue)			6.09	
1396	735296	( Committee Colon Vis. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)				
			6	0	6.09	
1397		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1397	13666					
		33,04 (Breast, High Met vs. Breast, Non-Met)	1	9		9.22
398	732712					
		5,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		6,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	3.02
399	3765			<del>-  </del>		
			<del>-   -  </del>	-+		
	c	1,02 (Colon, High Met vs. Colon, Low Met)	19	6	2.02	
400	185596		17	<u> </u>	2.92	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.
1401	1943					
		03,04 (Breast, High Met vs. Breast, Non-Met)		<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	4	3.9	+
	<u> </u>	01,02 (Colon, High Met vs. Colon, Low Met)	9	19 29	<del> </del>	1 3
1402	448193	, because one, see with the		1 29		3.4
		16.17 (Color Turner Time Color No.				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.08	+_
1403	1793	1932 (Torrian Colon Vs. Colon Turnor Hasue)	1	10		9.4
		01 02 (Colon High Materia Colon Line)				
		01,02 (Colon, High Met vs. Colon, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met)	13	27		2.2
1404	2475	(Breast, fight wet vs. Breast, Non-Met)	35	13	2.63	+-
		03.04/D				上
1405	730866	03,04 (Breast, High Met vs. Breast, Non-Met)	5	35		7.1
				-		+
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	8	0	8.12	
1406	730389	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.5
				$\vdash$		$\vdash$
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.5
1407	641884	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1408	463487	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7 11	-
1409	5156	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
1410	728408	01,02 (Colon, High Met vs. Colon, Low Met)	1	8		8.67
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	12		11.3
1411	73812	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	12	0	12.19	
1412	1662	03,04 (Breast, High Met vs Breast, Non-Met)	0	12		12.3
				$\dashv$		_
		01,02 (Colon, High Met vs. Colon, Low Met)	7	34		5.27
1413	736556	03,04 (Breast, High Met vs. Breast, Non-Met)	31	5	6.05	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.12	
1414	5240	(A Colonia Colon vs. Colon Tumor Tissue)	0	8		7.57
		08,09 (Lung, High Met vs. Lung, Low Met)	18	10	2.52	

Table 5

Table SEQ	CLST	T. D D.				· · · · ·
SEQ	CLSI		A	B	A/B	B/A
1415	(104	03,04 (Breast, High Met vs. Breast, Non-Met)	12	2	5.85	
1415	6184					
	<del> </del>	01.02/0.1 17.1 17.		_		
1416	446404	01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.67
1410	440404			<u> </u>		
	<del>                                     </del>	19 10 OL 1 C 1 . T' C T		-		<u> </u>
		18,19 (Normal Colon Tissue vs. Colon Tumor)	23	0	26.28	<u> </u>
<b></b> -		15,17 (Normal Colon Tissue vs. Colon Metastasis)	73	0	78.36	<u> </u>
	<b>-</b>	18,20 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	23	0	19.66	<u> </u>
1417	646825	13,10 (Normal Colon vs. Colon Tumor Tissue)	73	1	77.16	
	040023					-
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)		+-		
1418	734929	15,17 (Normal Coloir Tissue vs. Coloir Metastasis)	10	0	10.73	
1110	131727			₩	<del> </del>	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)		+_		1.0
	-	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.68
1419	648851	13,17 (Colon Tullor Fissue vs. Colon Metastasis)	6	10	6.09	<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		-	( 11	
1420	640135	( total Colon rissac vs. Colon Metastasis)	6	0	6.44	
				├──		-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.00	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	6.09 8.59	<del></del>
1421	7443	( Section Wedge 2015)		1	6.39	<del> </del>
				<u> </u>		
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	0	8.78	
1422	454050		<del>-   ´</del> -	<u> </u>	0.76	
				<del>                                     </del>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
1423	3765			<u> </u>	0.57	
		01,02 (Colon, High Met vs. Colon, Low Met)	19	6	2.92	
1424	648320					
				-		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1425	451269					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	4	3.49	
1.1-1		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	3	4.58	
1426	535208					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1427	720117	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1427	728115					
		151601 101				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1428	5240	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1428	5240					
		02 04 (Decent Health)				
		03,04 (Breast, High Met vs. Breast, Non-Met)	12	2	5 85	
1429	909	08,09 (Lung, High Met vs. Lung, Low Met)	18	10	2.52	
1.27	203				ł	1

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
	-	08,09 (Lung, High Met vs. Lung, Low Met)		+-	11.00	
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	34 54		2.93	+
1430	447697			110	2.93	
				<b>-</b>		1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	11		5.2
1431	447737					
	<del>                                      </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)				
1432	651100	13,10 (Normal Colon vs. Colon Turnor Hissue)	19	6	3.35	
				+	<del>                                     </del>	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	+
1433	735477			$T^-$		T
	<u> </u>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1434	3774	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	7	0	7.11	
1757	3174			┼		<u> </u>
···		01,02 (Colon, High Met vs. Colon, Low Met)	1	12		13.01
1435	646146		<del>-   -</del>	12		13.01
				1		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1436	642021	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1430	643931					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<u>                                     </u>
1437	463487	, ( Section inclusions)		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	7.51	
				<del>                                     </del>		<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	<u> </u>
1438	650097					
		16.17 (Colon Turner Tierre C.1. Maria				
1439	554469	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
				-		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	24	3	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	24	8	3.17	
1440	476223					
		15 17 01 1 0 1 7				
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	7	0	7.51	
1441	8738	23,24 (Normal Lung Tissue vs. Eung Tumor Tissue)	2	19		9.4
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
1442	403978					
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	23	50		2.15
1443	185539	15,16 (Normal Colon vs. Colon Tumor Tissue)	7	19		2.57
1173	103339					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	10		10.25
1444	451811		<del>-   '  </del>	10		10.25
				$\dashv$		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
1445	140731					

## Table 5

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Table						
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
	<del>                                     </del>					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1446	734582					
ļ	<del> </del>	151601101				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1447	463487	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6 09	<u> </u>
1117	403407			-		
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)		+-	0.50	
1448	558719	( Comme Colon Tibode 13. Colon Metastasis)	8	1	8.59	
				+-	_	<del></del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	18	6	3.17	+
1449	21669		- 1.0	+-	1 3.17	<del>                                     </del>
			·····	1		<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1450	470462					1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1451	3316					
		1610/01				
1452	552720	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1432	553728			_		
ļ		16 17 (Colon Turnor Tiesus va Colon Maria		╄┈	ļ	
1453	736014	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
	7,00017			╀—		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	-		5.60
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	6	6.09	5.68
1454	237288		<del>-   `</del>	<del>                                     </del>	0.09	<del> </del>
				<del> </del>		\ <del></del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1455	11141					
1456		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1456	556421					
		15 16 (Normal Colon or Colon T				
1457	549435	15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	
	3 13 130			_		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	2	7.4	
1458	448927	· (	14		7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
1459	379105					2.57
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
1460	552614					
1461		6,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1461	470602					
	<del>-  </del> .	51/01, 101				
		5,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	
		6,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	Т
1462	557039			+-	100	+
				+	<del></del>	+
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	16		+
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11		5.81	+
1463	549864					$\dagger$
	<del>                                     </del>	16.17 (Color T T				I
1464	449836	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	<b>_</b>	$\bot$
				<del></del>		+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	+
1465	554812					1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		8	-	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8	<del>                                     </del>	+
1466	3316			┪		+
		16.17 (Color Trans. T' C. 1. 24				
1467	649852	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		+
				╁	+	+
1160		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	T
1468	453592					I
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	6	3.35	╀
1469	455096			1	3.33	╁
	<del></del>					
-+	<del></del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9
1470	446199	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	10		4
				╁┈		├
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7
1471	558427					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	4	15		-
1472	450255	Tamer Hosacy		13		3
		16.17.60.1				
1473	452026	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5
				-		-
1474	27.1071	15,16 (Normal Colon vs. Colon Tumor Tissue)	35	14	2 64	
1474	374971					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.
		08,09 (Lung, High Met vs. Lung, Low Met)	0	16		11
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.
1475	446404					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	72			
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	73	0	78.36 19.66	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	73	1	77.16	
1476		18,19 (Normal Colon Tissue vs. Colon Tumor)	23	0	26.28	
1476	549591					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		- 1		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7 2	24		3.3

16,17 (Colon Tumor Tissue vs. Colon Metastasis)
15,17 (Normal Colon Tissue vs. Colon Metastasis)

Table 5
Page 95 of 155

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	T
1477	640135					+
				+	1	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	+-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	十
1478	646248					土
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		1	( 00	L
1479	639705	13317 (Colon Turnor Tissue vs. Colon Metastasis)	6	0	6.09	+
				+		╁
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	+
1.100	10200	15,16 (Normal Colon vs. Colon Tumor Tissue)	10	0	10.57	
1480	483084					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	+
1481	464029					+
		16.17 (Colon Tumor Tissue va Colon Maria				$oxed{\Box}$
1482	428005	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		1 8
				$\vdash$	<del>                                     </del>	╁╴
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		1 7
1400	01155	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7
1483	91178					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		<del>  6</del>
1484	550571		- i ·	<del>  '</del>		+ 0
						1
1405		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5
1485	735028					_
	<del></del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	+
1486	559409					
-		16,17 (Colon Tumor Tissue vs. Colon Metastasis)				<u> </u>
1487	551172	1031 (Colon Famor Florace vs. Colon Micrasiasis)	0	6		5.
		Mark Olivers				
1488	648872	15,16 (Normal Colon vs. Colon Turnor Tissue)	1	10		9.
	2.0012					$\vdash$
$ \top$		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1400	14645	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1489	446404					
		18,19 (Normal Colon Tissue vs. Colon Tumor)	23	0	26.28	_
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	23	0	19.66	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	73	1	77.16	
1400		15,17 (Normal Colon Tissue vs. Colon Metastasis)	73	0	78.36	
1490	734063			$\neg$		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	7.11	6.0
491	467991					
- 1						

Table 5

SEQ	CLST		A	В	A/B	
1492	454050					
	<del>                                     </del>	15.17.01			<u> </u>	
1493	734646	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	1
1493	/34040			+	<del> </del>	╀
		15,16 (Normal Colon vs. Colon Tumor Tissue)		14	<del> </del>	+
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	14	14	14.22	+
1494	450192	, (	17	Ť	17.22	╁
				1	<del>                                     </del>	+
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		T
1495	403978					T
	L	23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	23	50	ļ	_
1496	734209	15,16 (Normal Colon vs. Colon Tumor Tissue)	7	19		$\bot$
1470	734209			<del> </del>	<del>                                     </del>	╀
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	+
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9	9.14	╁
1497	14805			† ´		十
				<b>†</b>		1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	10		
1498	230995					$oxed{\Box}$
		151/ OL 10.1 G. I. T T.			<u> </u>	L
1499	120049	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		Ľ
1400	120049					╄
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	┢
1500	642142		- 1	<u> </u>	3.37	H
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
1501	402070	15,16 (Normal Colon vs Colon Tumor Tissue)	2	11		L.
1501	403978					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	19		_
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	23	50		2
1502	386543			50		É
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	23		7
1502	250105	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	23		7
1503	379105					<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		<del>  -</del>
1504	450255			°		7
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5
1505	730143					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6
1506	734209	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
.500	127403					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	0.
1507	401553		1	<del>-</del> +	Z T	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		1517 Olympic Ch. Ti			0.50	-
1508	72979	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1500	,2,,,	100		<del>                                     </del>		
		18,19 (Normal Colon Tissue vs. Colon Tumor)	1	36		31.5
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	36	4	6.73	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	46	311		6.3
		08,09 (Lung, High Met vs. Lung, Low Met)	18	0	25.15	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	46	193		3.97
1509	726307			ļ		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6		6.00	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	6.09	5.68
1510	230995	15,10 (Normal Colon vs. Colon Tunior Tissue)		l °		3.08
				-		<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1511	3524					
		01,02 (Colon, High Met vs. Colon, Low Met)	21	6	3.23	
1512	8112					
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	0	7.91	
1513	5240	05,04 (Bleast, High Met Vs. Bleast, Non-Met)		0	7.81	-
1313	3210					
		08,09 (Lung, High Met vs. Lung, Low Met)	18	10	2.52	
		03,04 (Breast, High Met vs. Breast, Non-Met)	12	2	5.85	
1514	447326					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1515	2676					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	17		9.22
1516	736701	01,02 (Colon, Tright Wict vs. Colon, Low Mict)		17		9.22
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	10	0	10.1	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1517	736701					
		1616 OL - 10.1 - 0.1 T - T				5.60
		15,16 (Normal Colon vs. Colon Tumor Tissue) 23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	10	6	10.1	5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1518	8371	100 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Table 1 To Tabl	Ť		0.07	
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1520	185542					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	10		10.25
1521	448046					
$\dashv$		15,17 (Normal Colon Tissue vs. Colon Metastasis)		12		11 10
-		16,17 (Normal Colon Tissue vs. Colon Metastasis)	3	12		3.94
1522	185422	(Colon Admir Albado Fai Colon Metastasis)	-	-14		3.34
			<del>  </del>	-		

Table 5

SEQ	CLST	Library Pair A,B	A	B	A/B	
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	32		
1523	650448					Т
						$\perp$
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1524	5753			1		L
		01.00 (0.1 )				┸
1526	1644	01,02 (Colon, High Met vs. Colon, Low Met)	0	10		L
1526	1644			<u> </u>	ļ	$\perp$
		02.04 (Proced High Mayor B. (N. N.)		4	<u> </u>	╄
		03,04 (Breast, High Met vs. Breast, Non-Met)	27	57		┸
1527	4453	01,02 (Colon, High Met vs. Colon, Low Met)	11	33		╀
1327	4400			—		+-
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	2	1.00	+
1528	454152	object (Steady Might Met vs. Diedst, Mon-Met)	10	+-	4.88	+-
				┼	<del></del>	+
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3.88	十
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	1	11.81	+-
1529	9913					╁
				T		╈
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	T
1530	1350					
		01,02 (Colon, High Met vs. Colon, Low Met)	3	44		1
1531	188			<u> </u>		
		02.04 (D				L
		03,04 (Breast, High Met vs. Breast, Non-Met)	129	309		2
1532	4471	21,22 (Normal Prostate vs. Prostate Cancer)	71	166		2
1332	77/1					_
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	H
1533	2622	, ( State of the Broads, 1 on 11cc)	- 10		9.70	-
						┢
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	31		4
1534	185465			-		<del> </del>
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	25	9	2.81	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14		_ 3
1535		03,04 (Breast, High Met vs. Breast, Non-Met)	2	14	•	7
1535	19205					
		02.04/David H'd M.				
1536	185635	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.
1000	103033					
		03,04 (Breast, High Met vs. Breast, Non-Met)		<del>-</del>		
1537	5289	(Siender, Angli Mott vs. Dicast, Noll-Wiet)	0	7		7.
		03,04 (Breast, High Met vs. Breast, Non-Met)	19	2	9.27	
1538	779		19	-	7.41	
			<del>-    </del>			
		01,02 (Colon, High Met vs. Colon, Low Met)	27	54		2.
		03,04 (Breast, High Met vs. Breast, Non-Met)	60	22	2.66	
539	779			-+		

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	
		01,02 (Colon, High Met vs. Colon, Low Met)	27	54		
		03,04 (Breast, High Met vs. Breast, Non-Met)	60	22	2.66	+
1540	5289			+-		$\dagger$
						1
		03,04 (Breast, High Met vs. Breast, Non-Met)	19	2	9.27	1
1541	456808					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	42		
1542	546642	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	42	1	42.65	L
1543	546642			-		1_
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		<del>                                     </del>	11.1	╄-
1544	649732	10,17 (Coloii Tullioi Tissue vs. Coloii Metastasis)	11	0	11.17	┼-
	019732					┼
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	╀
1545	5240		<del></del>	<del>                                     </del>	7.11	╫
						1
		03,04 (Breast, High Met vs. Breast, Non-Met)	12	2	5.85	十
		08,09 (Lung, High Met vs. Lung, Low Met)	18	10	2.52	$\vdash$
1546	448046					
						T
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	12		11
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	12		3
1547	650476					
		15 15 07 15 15 15				
1548	379341	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1340	379341					
		08,09 (Lung, High Met vs. Lung, Low Met)	2	21		_
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	21 14		4.
1549	401849			17		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	15		14
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1550	11452					
1551		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6
1551	185417					
		08 09 (Lung High Motors Lung Lung L				
-+	<del></del>	08,09 (Lung, High Met vs. Lung, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met)	8	56		5.0
1552	4471	OS, OT (Diedst, Trigil Met Vs. Diedst, Non-Met)	4	32		8
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	
1553	2557		10		9.70	
				-+		
		01,02 (Colon, High Met vs. Colon, Low Met)	23	8	2.65	
1554	3656			<del>-                                    </del>	2.03	
				$\dashv$		
		01,02 (Colon, High Met vs. Colon, Low Met)	2	12		6.:
1555	2327			_		
						-
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	19		2.4
1	į.	08,09 (Lung, High Met vs. Lung, Low Met)	10	2	6.99	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	T
1556	449026			1	1.72	+
				+-	<del></del>	+
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	╫
1557	730227			+-	3.57	╫
						+
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	+
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.05	┿
1558	650864		· ·	╁	<del>                                     </del>	+
				+-		╁
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	十
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	╁
1559	530774					╁
						十
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	5	16		$\top$
1560	395341					1
		151601 101				Γ
1561	557007	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		
1561	557906		_			
		15 16 (Normal Colon va Colon Transition				
1562	452531	15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	上
1502	432331			<u> </u>		1
		15,16 (Normal Colon vs. Colon Tumor Tissue)				╄
1563	559057	copie (Normal Colon V3. Colon Tunior Tissue)	2	11		⊢
						╀
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6
1564	448046		v	<del>                                     </del>		1
				$\vdash$		<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	12		1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	12		3
1565	553547					
1566		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5
1566	4636					
_		02 04 (Dreset History D. 1977)				
1567	455601	03,04 (Breast, High Met vs. Breast, Non-Met)	3	15		5.
-	100001					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.72	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	10.73 8.12	
1568	172013		-   0	-	0.12	
				- +		
		5,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1569	552597			$\neg$		
		5,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.:
1570	446531					
		(17/01 m m				
.571		6,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
311	639352			_		
_	1	5 17 (Normal Colon Treme are Colon Me				
-+		5,17 (Normal Colon Tissue vs. Colon Metastasis) 5,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.51	
	642604	2,10 (1101111a) COIOII vs. COIOII Tumor Hissue)	7	0	7.4	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	<u> </u>	10.72	ļ
	-	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.73	-
1573	558534			<u> </u>	1312	<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1574	556421			<u> </u>		
<b></b> .		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	<del> </del>	12.69	-
1575	735477	13,10 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	
	,,,,,,					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1576	640703					
		1617 (6.)				<u> </u>
1577	643878	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.16	-
1377	043878					<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1578	557797					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	16		5.25
1579	557200					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	10		0.05
		15,17 (Colon Tuthor Tissue vs. Colon Metastasis)	1	10		9.85
1580	729531			1.0		7.52
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1501	53.4554	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1581	734554					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	3.00
1582	418008					
1.500		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1583	558614					
	<del></del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
1584	452245	25,17 (Colon Tonor Toole V. Colon Metastasis)	1			0.00
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	12		11.18
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	12		11.82
1585	449891					
		15 16 Olympia Color Color Trans Trans				
1587	6162	15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
-207	0.02					
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	0	12.68	
1588	6162					
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	0	12.68	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		100 00 (I W 1 W 1 W 1 W 1		-		
	<del>                                     </del>	08,09 (Lung, High Met vs. Lung, Low Met) 01,02 (Colon, High Met vs. Colon, Low Met)	15	27	13.84	6 44
1590	3926	01,02 (Colon, Tingh Met vs. Colon, Low Met)		<u> </u>	13.04	
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
1591	185693					
1592	641683	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
1392	041083			-		<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
1593	11351					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1594	650864					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
1595	460445			Ť	00	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1596	447669					
		1516 01 10 1 10 1				
1597	227936	15,16 (Normal Colon vs. Colon Tumor Tissue)	4	16		3.78
1377	227750					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
1598	639459					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
1599	650195	15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	
1399	030193	V - 10 - 10 - 10 - 10 - 10 - 10 - 10 - 1				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1600	734793					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
1601	540787	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
1001	340707					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1602	400654					
1602	721467	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7 88
1603	731467					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	2.00
1604	4045					
1604	4045	10,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	9

Table 5

SEQ	CLST	Library Pair A,B	A	B	A/B	B
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	
		01,02 (Colon, High Met vs. Colon, Low Met)	2	11		5
1605	447669					
		15,16 (Normal Colon vs. Colon Tumor Tissue)		16		<u> </u>
1606	11351	13,10 (Normal Colon vs. Colon Tumor Tissue)	4	16	<del> </del>	1 3
1000	11331			1		╫
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		$\vdash$
1607	648931					1
1600	72.770.6	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
1608	726786					╂
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	╂
		15,16 (Normal Colon vs. Colon Tumor Tissue)		8	0.12	7
1609	4508	, (		Ť		† <i>'</i>
1616	41.70.77	01,02 (Colon, High Met vs. Colon, Low Met)	1	12		1.
1610	415058			<u> </u>		_
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	-		<u> </u>
1611	450633	10,17 (Colon Tunior Fissue vs. Colon Metastasis)		6		5
		15,16 (Normal Colon vs. Colon Tumor Tissue)	34	13	2 76	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	34	7	5.21	
1612	736955					
		15 16 (Normal Calar va Calar Trans Trans		10		L.
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	13	13.2	1
1613	729851	10,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	0	13.2	-
				-		H
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.
1614	2512					
		03 04 (Proper Uick Matrix Proper Nov. Mat)				
1615	452704	03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	1	10.57	
1616	4589					
	*					
1617	4727	01,02 (Colon, High Met vs. Colon, Low Met)	14	2	6.46	
1617	4727					
		03,04 (Breast, High Met vs. Breast, Non-Met)	19	0	18.54	
1618	454380			<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	14		4.
619	553912					
$\dashv$		16,17 (Colon Tumor Tissue vs. Colon Metastasis)				
		15,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.8
			1 1/ 1	/		D.

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	13		12.3
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	2	6.6	
1621	448193					
	L		<del> </del>			ļ
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	2	5.08	<b> </b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10		9.46
1622	549591				<u> </u>	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	24		11 10
		16,17 (Normal Colon Tissue vs. Colon Metastasis)	7	24		3.38
1623	448511	10,17 (Colon rumor rissue vs. Colon ricustasis)	+	24		3.36
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	26		8.07
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	26		6.4
1624	335		ļ			<u> </u>
		12 14 GEORGE WALLIAMS TO A MEGET WALLIAMS TO	<del>                                     </del>	15		1.02
		13,14 (bFGF Treated HMVEC vs. VEGF-Treated HMVEC)	3	15		4.92
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	38		3
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	41		3.18
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	29	5.43	25.38
		19,20 (Colon Tumor Tissue vs. Colon Metastasis) 12,14 (Untreated HMVEC vs. VEGF-Treated HMVEC) (Untreated	29	4	5.42	
		HMVEC vs. VEGF-Treated HMVEC)	1	15		14.69
1625	561382					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	2	6.34	<b> </b>
1626	3447					
		08,09 (Lung, High Met vs. Lung, Low Met)	0	13		9.3
		01,02 (Colon, High Met vs. Colon, Low Met)	2	16		8.67
1627	639896		1			
1.000	1252	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
1628	1353	<del> </del>	+			<b> </b> -
		01,02 (Colon, High Met vs. Colon, Low Met)	39	13	2.77	
		03,04 (Breast, High Met vs. Breast, Non-Met)	37	17	2.12	
1629	3031	ossis (States) anguitation of States (Vol. 1924)	1	1		
		01,02 (Colon, High Met vs. Colon, Low Met)	7	18		2.79
1630	557928					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	6		5.91
1631	4727	10,17 (Colon Tumor Tissue vs. Colon Metastasis)	1-	-		3.91
			1			
		03,04 (Breast, High Met vs. Breast, Non-Met)	19	0	18.54	
1632	4046					
			1			2.70
1633	10882	01,02 (Colon, High Met vs. Colon, Low Met)	4	14		3.79
1033	10002		<del>{</del> {			
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	20		3.42
1634	646283					
			11			

Table 5 Page 105 of 155

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	<b>B</b> /2
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9.51	
1635	646283					ļ
		15,16 (Normal Colon vs. Colon Turnor Tissue)	9	0	9.51	
1626	120516	15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	-
1636	139516					-
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.1
1637	6184	oso ( (Broad, 11g. 11e to Broad, 1 or 11e)				
		01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.6
1638	6184					<u> </u>
						_
1620	454652	01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.6
1639	454653					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	5	17		3.3
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	17		5.2
1640	3309					
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	0	8.78	
		08,09 (Lung, High Met vs. Lung, Low Met)	10	38		2.1
1641	1037					
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	22		5.0
1642	450665	05,04 (Breast, High Met vs. Breast, Non-Met)				
	100000					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
1643	726307					
						<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	-
1644	447669	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.0
1044	447009					_
		15,16 (Normal Colon vs. Colon Tumor Tissue)	4	16		3.1
1645	639651				· · · · · ·	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1646	736860					_
		16.17 (Colon Turnor Tienna ve Colon Metadoria)		0	6.00	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	6.09	5.6
1647	553705	15,10 (110111121 COIOH VS. COIOH TUHIOI TISSUE)		9		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	0	12.68	
1648	451375					
				$oxed{oxed}$		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	11		5.
1649	204862					<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	$\vdash$

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1651	447539					
1037	11,000					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	<u> </u>
1650	455006	13,17 (Normal Colon Tissue vs. Colon Metastasis)		$\stackrel{}{-}$	0.07	<del>                                     </del>
1652	455096				No. today	-
		(1 M · · · · )	2	10		4.92
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		10		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32
1654	449142					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	1	9.14	
1655	557401				_	ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	
1656	418763					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
1657	17649					
	A.	03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
1658	2078	(				
1050	20,0					
		03,04 (Breast, High Met vs. Breast, Non-Met)	11	2	5.37	
1650	640370	103,04 (Bleast, High Met 43. Bleast, 1101 Met)	<del></del>			<del> </del>
1659	040370			-		
		16 17 (Color Towns Tissue as Color Metestasis)	7	0	7.11	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.44	
1660	440060	15,17 (Normal Colon Tissue vs. Colon Metastasis)			0.44	
1660	449269			$\vdash$		
		Is a second of the second of t	26	7	3.99	<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)				<b></b>
	l	15,16 (Normal Colon vs. Colon Tumor Tissue)	26	5	5.5	<b>-</b>
1661	639029					├
						<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<u> </u>
1662	448677					<u> </u>
						<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	1	11.63	
1663	349					
		01,02 (Colon, High Met vs. Colon, Low Met)	69	138		2.1
		03,04 (Breast, High Met vs. Breast, Non-Met)	77	1	75.13	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.5
1664	447494					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	26	8	3.44	
1665	551433					
	<b> </b>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.8
1666	414739	10,17 (Colon Tunior Tibbue 10, Colon Promomono)				T
1000	717/39					
	1	15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	14		6.5
			2	14		6.8
	1	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		14		1 0.8

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	ļ
1668	640525					<u> </u>
		15 17 Oleman Calan Tirana na Calan Matastasia)	7	0	7.51	
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Turnor Tissue)	7	0	7.31	
1669	233108	15,16 (Normal Colon vs. Colon Tuntor Tissue)		0	7.7	
1009	233100					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	0	11.63	
1670	643594					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1671	1642					
		03,04 (Breast, High Met vs. Breast, Non-Met)	28	5	5.46	
1672	643804					<u> </u>
		15 17 (Namual Calan Tragua ya Calan Matastasia)	6	0	6.44	
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1673	449701	13,10 (Normal Colon vs. Colon Tunior Tissue)			0.54	
1075	449701					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	17		16.08
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	17	1	17.26	
1674	185695					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
1675	555830					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	2	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	16	2	8.59	<u> </u>
1676	227936					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
1677	1609	15,16 (Normal Colon vs. Colon Turnor Hissue)		0		1.51
1077	1007					E
		01,02 (Colon, High Met vs. Colon, Low Met)	3	58		20.96
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	3	5.37	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	14	3	4.74	
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	2	4.88	
1678	643938					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1679	3656					
		01.02 (Colon High Meters Colon Law Max)		12		6.5
1600	16576	01,02 (Colon, High Met vs. Colon, Low Met)	2	12		6.5
1680	16576					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2
		object (Dieust, 111811 Free vs. Dieust, 11011-11101)		۲		3.2
1681	9784	ł .	l l			

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
<del>-</del>		03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
1682	2557					1
		01,02 (Colon, High Met vs. Colon, Low Met)	23	8	2.65	1
1683	4620					
						l
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	i
1684	43642					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1685	555103					
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	0	6		5.94
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1686	643341					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1687	185531			l		
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	9		9.22
		08,09 (Lung, High Met vs. Lung, Low Met)	0	13		9.3
1688	4045					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	11		5.96
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	
1689	400258					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1690	96618					
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	13		3 33
1691	646060					<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	0	13.95	
1692	5((5	15,16 (Normal Colon vs. Colon Tumor Tissue)	13	3	4.58	
1092	5665					
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	0	12.68	
1693	149265	05,04 (Bleast, Figil Met Vs. Bleast, Non-Met)	13	0	12.08	
1023	149203					
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	16		16.4
1694	727314	ve jo . (2. 1000) right tract to Diodog Hole Hotel				10.4
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	3.32
1695	736349					
			<del></del>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1696	648931	<u> </u>				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
1697	553881					

Table 5

Table :						
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	4	3.96	
1698	7444					
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	0	8.78	
1699	150					
				-		
	<del> </del>	03,04 (Breast, High Met vs. Breast, Non-Met)	51	24	2.07	ļ — — —
		08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	
1700	2889	08,09 (Lung, High Met vs. Lung, Low Met)	<del>-   -</del> -	<del>                                     </del>	0.55	
1700	2009	<del>                                     </del>		├─	<del>                                     </del>	<del> </del>
		01 02 (Colon High Metays Colon Levy Met)		19	<b></b>	257
1701	720670	01,02 (Colon, High Met vs. Colon, Low Met)	8	19		2.57
1701	730670			├—	<b> </b>	
				<del>  </del>		<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	ļ	5.68
1702	560984	<u> </u>		<u> </u>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1703	453708			1		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	19		18.71
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	19		17.7
1704	48977					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
1707	97507			1		
	7,150	<del> </del>		-		
	<del>                                     </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	1	9		8.51
1708	735966	15,10 (Normal Colon Vs. Colon Tattion 1135ac)	<del></del>	l-		0.51
1700	133700			<del> </del>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16.17 (Colon Turnor Tissue vs. Colon Metastasis)	6	0	6.09	3.08
				0	7.07	
1700	25	23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	7	"	7.07	
1709	35	<del> </del>		<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
		03,04 (Breast, High Met vs. Breast, Non-Met)	386	1967		5.22
		08,09 (Lung, High Met vs. Lung, Low Met)	868	11	110.27	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	14		6.62
1710	650195					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1711	639705					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	0	10.57	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
1712	185465					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	14		3.45
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	14		7.17
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	25	9	2.81	
		1 , ( ) Dung Tuntor Tibuto)		ليئيا	2.01	

Table 5

Table:						
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1713	378525					1
				T		
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	9	1	9.14	
1714	2889					
		01,02 (Colon, High Met vs. Colon, Low Met)	8	19		2.57
1715	557686					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1716	735786			1		
				1		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	<del> </del>	6.62
		16,17 (Colon Tumor Tissue vs Colon Metastasis)	7	0	7.11	<del> </del>
1717	455145		<del></del>	<del>                                     </del>		-
				<del>                                     </del>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<del>                                     </del>
1718	639667	10,17 (Yournal Colon 1,0000 10 Colon Monacasta)	<del>+</del> -	<del>                                     </del>	0.11	<del> </del>
	333007	<del> </del>		1		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
1719	446913	10,17 (Colon ranio rissae vs. Colon racastasis)	<del>+-</del> -	<del>                                     </del>	2.14	
1715	170713	<del></del>		<del> </del>		<del> </del>
	<del> </del>	18,19 (Normal Colon Tissue vs. Colon Turnor)	10	0	11.43	<del> </del>
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	63	0	67.62	
	<del> </del> -	18,20 (Normal Colon Tissue vs. Colon Metastasis)	10	0	8.55	
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	63	1	66.59	
1720	402494	13,16 (Normal Colon Vs. Colon Tullior Tissue)		1	00.39	
1720	402494			<del> </del>		<b></b> -
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	3	13	<del> </del>	4.1
	<del> </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	3	4.4	4.1
1721	734256	10,17 (Coloit Tullioi Tissue vs. Coloii Metastasis)	13	+-	4.4	
1/21	734230			├	<del></del>	
		15 16 Oleman Colon va Colon Transa Tinava	<del></del>	7	<del>[</del>	((2)
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7		7.11	6.62
1722	734256	10,17 (Colon Turnor Hissae vs. Colon Metastasis)	<del></del>	0	7.11	
1722	734230					
		16 17 (Colon Turner Tisque va Colon Meteotoria)	<del></del>	-	7.11	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	$\frac{7}{2}$	0	7.11	
1723	550262	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1723	559362					
		1/ 17/0 1 T T O 1 1/ 1/ 1	<del></del>	-		
1704	(20(6)	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	<u>l</u>	8		7.88
1724	639651		<del></del>			
	ļ	15 17 OL 1 C - 1 - T'				<u></u>
1725	410774	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1725	419774	<del></del>				<u> </u>
1706	555010	15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
1726	555318	<del> </del>				
		1617 (6.1 )				
155	1105 = 5	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	12		5.91
1727	449956					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	
1728	558427					

Table 5

Table 5						
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	4	15		3.55
1729	7531					
		01,02 (Colon, High Met vs. Colon, Low Met)	1	8		8.67
1730	446514					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	
1731	456808				· · · · · · · · · · · · · · · · · · ·	
				<u> </u>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	42	1	42.65	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	42		5.68
1732	447035			<u> </u>		
	1111111					
	<del>                                     </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	
1733	446913	10,11 (Colon Tamor Tibbae 13. Colon Metababis)		<del>                                     </del>	0.12	
1,33	410713		<del></del>			
	<del> </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	63	1	66.59	<del>                                     </del>
-	<b></b> -	15,17 (Normal Colon Tissue vs. Colon Metastasis)	63	0	67.62	<del> </del>
		18,19 (Normal Colon Tissue vs. Colon Tumor)	10	0	11.43	<b>-</b>
	<del></del>	18,20 (Normal Colon Tissue vs. Colon Metastasis)	10	0	8.55	
1734	446900	16,20 (Normal Colon Tissue vs. Colon Metastasis)	10	<del>ا</del> ّ	6.55	ļ
1754	440700			-		<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	11		5.42
		15,17 (Colon Tunior Fissue vs. Colon Metastasis)	$+\frac{2}{1}$	11		10.25
1735	504513	13,17 (Normal Colon Tissue vs. Colon Metastasts)		- ' '		10.23
1733	304313			_		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1736	380477	13,17 (Normal Colon Tissue vs. Colon Metastasis)		-	0.44	
1730	300477					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1738	8259	13,17 (Normal Colon Tissue vs. Colon Metasiasis)		۳	7.51	
1736	8239					
		01,02 (Colon, High Met vs. Colon, Low Met)	10	0	9.22	
			13	49	9.22	2.7
1720	9250	08,09 (Lung, High Met vs. Lung, Low Met)	13	49		2.7
1739	8259			_		
		01.02 (Colon High Mot ve Colon Levy Met)	10	0	9.22	
		01,02 (Colon, High Met vs. Colon, Low Met) 08,09 (Lung, High Met vs. Lung, Low Met)	13	49	9.22	2.7
1740	552968	08,09 (Lung, High Met vs. Lung, Low Met)	13	49		2.7
1740	332908			-		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.86
		15,17 (Colon Tumor Fissue vs. Colon Metastasis)	0	9		
1741	650045	13,17 (Normal Colon Tissue vs. Colon Metastasis)	<del> -</del>	- 9		8.38
1741	650845					
		16.17 (Color Turnor Tiggue va Color Material)	<del></del>		6.00	
1742	640504	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	- 6	0	6.09	
1742	648594	<del> </del>				
	<del></del>	15 17 (Normal Colon Tiesus v. Colon Material)			10.73	
17/0	(40.004	15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
1743	648594	<del> </del>				
		15 17 01 10 1 7 2 1 2 1			10.72	
1011	0707	15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
1744	2796					
		02.04 (Days Hall Mars Days N. 20.		-22		151
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	37		4.74

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Table 5	5					
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1745	5753					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	10		10.84
1746	734256					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1747	449580					
1747	117500					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	4	4.23	
1748	553705	15,10 (Norman Colon to Colon Tunior Tibeac)				
1740	333103					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	0	12.68	
1510	720670	15,16 (Normal Colon vs. Colon Tullor Tissue)	- 12	Ľ	12.00	
1749	730670		-			
			0	6		5.68
		15,16 (Normal Colon vs. Colon Tumor Tissue)			6.00	3.08
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1750	15035					
					0.46	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
1751	394436					
						<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<u> </u>
1752	726810					
						<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	<u> </u>
1753	352763					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	10		9.85
1754	3506					
<b></b>		03,04 (Breast, High Met vs. Breast, Non-Met)	1	10		10.25
1755	726377			i		
	120011					
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
	<del>                                     </del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1756	562111					
1,50						
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	13		64
<b> </b>	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	13		6.06
1757	404475	15,17 (Political Colon Pissas Vs. Colon Picsas Vs.		-		
1737	104473					
<u> </u>		16.17 (Colon Tumor Tissue vs. Colon Metastasis)	11	2	5.59	
<u> </u>	<del> </del>	15,17 (Colon Turnor Tissue vs. Colon Metastasis)	19	2	10.2	<u> </u>
1750	12024	13,17 (POLITICAL COLOR TISSUE VS. COLOR PACKASIASIS)	<del>- + ''</del>	+-	1	1
1758	13824			-	<del>                                     </del>	+
<u></u>	<del> </del>	02.04 (D 11:-1.14 (	6	0	5.85	+
	<b></b>	03,04 (Breast, High Met vs. Breast, Non-Met)	- 0	+ "	3.03	<del> </del>
1759	558222			₩-	<del> </del>	-
				+-	<del> </del>	5.01
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1760	2834			-		<del> </del>
1	1				<u> </u>	

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		01,02 (Colon, High Met vs. Colon, Low Met)	8	22		2.98
1761	453470					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1	12.68	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	1	12.88	
1762	558682	-,-,-				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	12		3.9
1763	641710					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1764	640221					
1,0.	0.000					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1765	559057					
1,03						
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.5
1766	551433					
.,	551.05					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.8
1767	5729					
	0,127					
		01,02 (Colon, High Met vs. Colon, Low Met)	1	10		10.
1768	352763					
1.00						
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	10		98
1769	375651					
1,05	3,3031					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
1770	644032					
1770	011002					
	l	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	124	0	125.92	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	124		16.
1771	185562					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.2
1772	736349					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.6
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1773	638870					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
1774	649719					
_						
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	ļ
1775	62016					
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
1776	2889					
_ , , , ,				<b>1</b>		
	<u> </u>	01,02 (Colon, High Met vs. Colon, Low Met)	8	19		2.5
1777	647135	, (,,,,,,,,,,,,,	<del></del>	1		T

Table 5

Table 5						1 - 1 - 1
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1778	8283					
						0.67
		01,02 (Colon, High Met vs. Colon, Low Met)	0	8		8.67
1779	732121					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1780	532307					
						6.00
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1781	6589					
						0.76
		01,02 (Colon, High Met vs. Colon, Low Met)	1	9		9.76
1782	554678					<del> </del>
			- 10	-	5 10	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	<del> </del>
1783	450410			<b>_</b>		<del> </del>
				7		6.89
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	<del>- '- '</del>		0.89
1784	643924					
		Large Colonia Colonia Torrio			6.34	
	<u> </u>	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.44	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	<u> </u>	0.44	<del>                                     </del>
1785	453719			<del>                                     </del>		
		16 17 (Color Transport Color Materials)	0	9		8.86
1706	451011	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	-			0.00
1786	451811			-		<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	2	5.37	<del>                                     </del>
1787	453059	13,17 (NOTHING COION TISSUE VS. COION IVICLASIASIS)			3.57	
1787	433039					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
1788	453457	13,10 (Nothal Colon Vs. Colon Tunior Tissue)				
1700	133137					
-		16.17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
1789	558454					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1790	417467					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1791	447850					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1792	557948					
	<u> </u>					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
1793	452685					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	5	3.17	
1794	446964			<u> </u>		

Table	:
SEQ	

Table 5								
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A		
		<u> </u>		<u> </u>				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32		
1795	550318							
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	3	5.28			
1796	407077							
<del>  </del>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88		
1797	650864	Toy. (cook rank hour to cook water by	<del></del>	<u> </u>				
1777	050001	<del> </del>		<del> </del>		<del> </del>		
<del> </del>		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	<del> </del>		
<del>   </del>				<del></del>	8.59	<del> </del>		
1700	(14501	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.39	<del> </del>		
1798	644721		-+			<del> </del>		
			<del>-  </del>					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<b> </b>		
1799	485431			ļ		<b>!</b>		
				<u> </u>		ļ		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68		
1800	651073			L				
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4			
1801	725811							
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68		
1802	645139							
	<del></del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	<del>                                     </del>		
1803	185478							
1005		<del> </del>	<del>-  </del>					
<del>  </del>		03,04 (Breast, High Met vs. Breast, Non-Met)	0	14		14.35		
1804	1441	05,01 (Diedst, 11igh Met 13. Diedst, 10in 11it)	— <del>      </del>	<u> </u>		1 4.55		
1004	1771		<del>-  </del>	<del> </del>		<del></del>		
<del>  </del>	<del></del>	01,02 (Colon, High Met vs. Colon, Low Met)	9	40		4.82		
		<del></del>	38	16	2.32	4.62		
1805	640005	03,04 (Breast, High Met vs. Breast, Non-Met)	- 30	10	2.32	<del></del>		
1803	040003		<del></del>			<del> </del>		
		161(0) 101 C1 T	<del>-  </del>		2.52	<b>}</b>		
<del>  </del>		15,16 (Normal Colon vs. Colon Tumor Tissue)	20	6	3.52	<del>                                     </del>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<b></b>		
<u> </u>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	20	0	21.47	<b>}</b>		
1806	728273							
						ļ		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62		
1807	185579							
L1		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2		
1808	724473							
	<del> </del>							
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09			
<del></del>		21,22 (Normal Prostate vs. Prostate Cancer)	5	16		3.25		

## Table 5

Table 5	CLST	Library Pair A,B	A	В	A/B	B/A
1809	559674	Indian y Kani Page		1	7102	10/21
1007	337074	<del>                                     </del>		<del> </del>		├
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7	<del></del>	6.52
1810	456026	15,11 (Voltilai Cololi 11,5500 VS. Cololi 130000055)	<del>-                                     </del>	<del>                                     </del>	<del></del>	0.52
1010	150020			1	<del> </del> -	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	24	4	6 34	<del>                                     </del>
<u> </u>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	24	8	3.22	<del> </del>
1811	549320					<b>†</b>
					<u> </u>	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1812	447338					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1813	560700					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5 91
1814	3070					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	L	5.91
1815	3070			<u> </u>	<u> </u>	<u> </u>
					<u> </u>	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1816	380477			ļ		
<u> </u>				<del> </del>		<b> </b>
1017	525040	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1817	735040	<del> </del>		<del>                                     </del>		
<b> </b>		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
	_ <del>_</del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	3.08
1818	378525	10,17 (Colon Funor Fissue vs. Colon Metastasis)	<del>+</del> -	├-	0.09	
1010	316323			1		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	1	9.14	<del>                                     </del>
1819	284586					
				-		<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1820	640276					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
1821	3344					
		01,02 (Colon, High Met vs. Colon, Low Met)	7	17		2.63
1822	555830					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	16	2	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	16	2	8.59	
1823	726307					<u> </u>
		151621 101 0 1 7	<del></del>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	( 00	5.68
1924	116	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	ļ ——
1824	416					<b> </b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	23		2.72
	<del></del>	21,22 (Normal Prostate vs. Prostate Cancer)	11	31		2.72
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	22		2.56
		20,1. (A TOTTIMI COTOTI FISSUE VS. COTOTI IVICIASIASIS)	<u> </u>	L		2,50

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1825	2543					
		01,02 (Colon, High Met vs. Colon, Low Met)	23	7	3.03	
1826	639352					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
1827	453592			L		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	6	3.35	<u> </u>
1828	450633					<u> </u>
				<u> </u>		ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	34	13	2.76	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	34	7	5.21	ļ
1829	448383			<u> </u>	ļ	<u> </u>
						<b> </b>
	110710	15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	1	13.95	<b></b> -
1830	648719			<b>-</b>		<b></b>
		15 17 01 - 10 1 T			0.66	<del>                                      </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	<del> </del>
1831	730655	15,16 (Normal Colon vs. Colon Tumor Tissue)	<del></del>	1	9.51	<del> </del>
1831	730033			├		<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<del> </del>
	<del></del>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.07	5.68
1832	141185	15,10 (Normal Colon Vs. Colon Tuntor Tissue)	<del></del>	-		3.08
1052	141103	<del> </del>				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	<del> </del>
1833	640498					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1834	9029					
		01,02 (Colon, High Met vs. Colon, Low Met)	7	0	6.46	
1835	559674					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1836	555734					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	10		9.32
4005	4040	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	10		9.85
1837	1943					<b> </b>
		02.04 (Day of W. 1 May - Day of W. 1 May)	<del></del>		2.0	
		03,04 (Breast, High Met vs. Breast, Non-Met)	16	4	3.9	2.40
		01,02 (Colon, High Met vs. Colon, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue)	9	29 19		3.49
1838	648320	15,10 (Normal Colon vs. Colon Tumor Tissue)	6	19		
1030	070320					<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<del> </del>
1839	558098		- <u>`</u> -	۲	0.57	<del> </del>
					L	<del> </del>
						1

Table 5

Table 5	CLST	Library Pair A,B	1 🛦	В	A/B	B/A
SEQ		Library Pair A,B	A	B	A/B	B/A
1840	468672	<del> </del>		├		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
		15.16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	7.66
1841	456596	15,10 (Normal Colon Vs. Colon Turnor Tissuc)	<del>+</del> -	₩,	0.54	<del></del>
1041	430370	<del></del>	<del>-</del>	-		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	12		11.82
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
1842	649722					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1843	550708					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
1844	643931					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	Ĺ
1845	726927				L	
				<u> </u>		L
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
1016	150010	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	11		10.41
1846	459012	<del> </del>	<del></del>			
		15 17 Olement Color Transcer Color Metadosis			<i>4 11</i>	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.44	
1847	397773	15,10 (Normal Colon vs. Colon Tunos Tissue)		-	0.54	
1047	391113					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	0	12.88	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
1848	450004					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	13		12.3
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	2	6.6	
1849	649732					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1850	553955					
1051	646200	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1851	646309			$\vdash$		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		0	6.09	
1852	402727	10,17 (Colon Tunior Tissue vs. Colon Metastasis)	<del>   ° -</del>	-	0.09	
1632	402727			-		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	5	17		3.17
1853	468736	1332 (1707) All Colon Hisac (6. Colon Household)				3.17
				$\vdash$		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1854	650422	, , , , , , , , , , , , , , , , , , , ,		┌┷┤		
			<del></del>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	18	6	3.17	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	18	0	19.32	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
1855	730533					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1856	726307					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.07	5.68
1857	450311					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	ļ
1858	450940					
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)		9		8.38
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.86
1859	726786	10,17 (Coloit Fullor 1134C vs. Coloit McCasasis)	<del> </del> -			0.00
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
1060	7624	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	ļ
1860	7634			-		<del> </del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
1861	230995					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1862	374770					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
1863	9275					
		03,04 (Breast, High Met vs. Breast, Non-Met)		6		615
1864	553860	object (Section 11) and the section of the section				
	<del></del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)		9		8.38
1865	452010	15,17 (Normal Cololi Tissue vs. Colon Metastasis)		9		8.38
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	3	4.4	
10//	(107(0	15,16 (Normal Colon vs. Colon Tumor Tissue)	3	13		4.1
1866	649560	<del> </del>		$\vdash$		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1867	452704					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	1	10.57	
1868	447594					
		18,19 (Normal Colon Tissue vs. Colon Tumor)	11	2	6.29	<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	20	J.27	3.11
1869	555444					
		16.17 (Color Turner Treese up Color Material)	-   22		<i>5.5</i> 0	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	22	22	5.59	10.41
1870	736556	15,10 (Normal Colon vs. Colon Tumor Tissue)	<del></del>	22		10.41

1884

1885

456764

641373

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8	0.12	7.57
1871	5289	15,10 (Normal Colon vs. Colon Tumor Tissue)				7.57
		03,04 (Breast, High Met vs. Breast, Non-Met)	19	2	9.27	
1872	732121					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1873	452567					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	14		4.6
		15,16 (Normal Colon vs. Colon Tumor Tissue)	21	3	7.4	
1874	551634					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.9
1875	644099					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1876	726788					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.6
1877	638802					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1878	646283					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	0	9.51	
1879	8403					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
1880	2224					
		03,04 (Breast, High Met vs. Breast, Non-Met)	44	8	5.37	
1881	650053					
	·	15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	1	10.57	
1882	380477					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1883	450867					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	

10

14

15

10.16

15.03

5.28

16,17 (Colon Tumor Tissue vs. Colon Metastasis)

15,17 (Normal Colon Tissue vs. Colon Metastasis)

15,16 (Normal Colon vs. Colon Tumor Tissue)

Table 5

Table 5						
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	15	0	16.1	
1886	555882					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.86
1887	644046					l
						<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
1888	447250					
		<u> </u>				
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6 09	
1889	456596					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	12		5.59
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	12		11.82
1890	2218					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	21		11.38
		03,04 (Breast, High Met vs. Breast, Non-Met)	27	8	3.29	
		08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	ļ
1891	446450					
						L
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3.88	
1892	640889					<u> </u>
						ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1893	530774			$\vdash$		<b></b>
			<del></del>			2.00
1004	(40062	15,17 (Normal Colon Tissue vs. Colon Metastasis)	5	16		2.98
1894	649062			<b> </b>		
		15 17 Olympia Color Times on Color Materials		$\vdash_{\sim}$	6.44	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1895	12808	15,16 (Normal Colon vs. Colon Tumor Tissue)		<b>├</b>	0.34	
1093	12008					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	<del> </del>
1896	468672	03,04 (Bleast, Trigit Wet vs. Bleast, Noti-Net)			3.63	<del></del>
1070	700072			$\vdash$		
		15,16 (Normal Colon vs Colon Tumor Tissue)	6	0	6.34	-
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	0	8		7.88
1897	650773					
						<b></b> -
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1898	732237					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1899	650773					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1900	550216					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1901	639189					

Table 5

Table 5	;					
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
1902	3447					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	16		8.67
		08,09 (Lung, High Met vs. Lung, Low Met)	0	13		9.3
1903	2012					
		03,04 (Breast, High Met vs. Breast, Non-Met)	20	58		2.97
		01,02 (Colon, High Met vs. Colon, Low Met)	13	29		2.42
1904	642876					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
1905	449690					
	<b>-</b>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	17		5.58
1906	451208					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	3	5.01	
1907	725811					
.,,,						T
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<b> </b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1908	1256	13,10 (Normal Colon to Colon Tames)				
1908	1230					
		08,09 (Lung, High Met vs. Lung, Low Met)	35	110	-	2.25
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	14	31		2.27
1909	446599	05,04 (Diedst, High Wet 15. Diedst, 1101 120)				<b>†</b>
1909	440377					1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	2	6.87	
1910	446537	13,10 (Normal Color to Color Lance Color)				
1710	440337					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
1911	726281	13,10 (Norman Colon Vs. Colon Tamor Tissue)				<del>                                     </del>
1911	720281					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	1
1912	11286	10,17 (Colon Tullior Tissue vs. Colon Metastasis)		-		+
1912	11200					+
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1012	556082	01,02 (Colon, 11igh Wet vs. Colon, Low Mety		Ť		1
1913	330082			$\vdash$		+
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	10		9.85
1014	07507	16,17 (Colon Turnor Tissue vs. Colon Metastasis)		10		1
1914	97507			<del> </del>		<del> </del>
	<u> </u>	15.16 Olympia Colon are Colon Turnor Tiggue)	1	9		8.51
1015	#25055	15,16 (Normal Colon vs. Colon Tumor Tissue)	- 1	"		+ 0.51
1915	535955			-		+
<del> </del>	<u> </u>	16.17 (Color Towns Times Color Messesia)	0	10		9.85
ļ	<del> </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		10		9.83
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32
1916	728251			-		1
	<u> </u>			<del> </del>	6.00	+
	<del>                                     </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6 09	5.00
Į .		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68

Table 5

Table 5								
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A		
1917	733849							
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62		
1918	447574							
		15,17 (Normal Colon Tissue vs. Colon Metastasıs)	7	0	7.51			
1919	7607							
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	3	4.65			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	4	3.44			
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59		
1920	644032							
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	124		16.76		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	124	0	125.92			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51			
1921	454087							
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	3	4.93			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	1	15.03			
1922	412364							
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44			
1923	535208							
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44			
1924	644609							
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51			
1925	645073							
1923								
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34			
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44			
1926	417467							
	<b></b>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52		
1927	554188							
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34			
1928	647185							
	1	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44			
1929	736679							
	1					T		
	<b> </b>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09			
1930	553547			i i		1		
	1 2233.7							
	<del>                                     </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91		
1931	641524							
	1			1				
	<u> </u>			<del></del>				

Table 5

Table 5						70.4
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1932	649717					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1933	451041					
						5.01
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1934	3483					
						7.22
		01,02 (Colon, High Met vs. Colon, Low Met)	3	20		7.23
1935	500959					
					0.46	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1936	500959					
					0.46	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
1937	697			$oxed{oxed}$		
						2.46
		03,04 (Breast, High Met vs. Breast, Non-Met)	30	72	4.00	2.46
		21,22 (Normal Prostate vs. Prostate Cancer)	10	2	4.92	
1938	736955					
					40.0	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	0	13.2	12.2
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	13		12.3
1939	554742					
Ĺ						5.01
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1940	642973					ļ
				<u> </u>	7.11	ļ
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
1941	449437			<del>                                     </del>		<del></del>
			14	1	4.93	ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	12	4.93	3.94
		16,17 (Colon Turnor Tissue vs. Colon Metastasis)	3	12		3.94
1942	467991			├		<del> </del>
<u></u>			7	0	7.4	<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)		H	7.4	<del>                                     </del>
1943	650204			┼		<del>                                     </del>
	<b> </b>	Legacian Color Materials	11	0	11.81	<del>                                     </del>
<b></b>	(12.55	15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	+ -	11.01	<del>                                     </del>
1944	640618			$\vdash$		
<u> </u>	ļ. —	15,17 (Normal Colon Tissue vs. Colon Metastasıs)	6	0	6.44	
1215	452265		<del>-   `</del>	<del>Ť</del>		
1945	452366					<del>                                     </del>
<u> </u>		15 17 Olympal Colon Ticque ve Colon Metastagia)	6	0	6.44	<del>                                     </del>
<u> </u>	<del> </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.34	
	60000	15,16 (Normal Colon vs. Colon Tumor Tissue)		+ -	0.34	+
1946	640276			+	<del> </del>	<del> </del>
	<del>                                     </del>	16.17 (Calon Tumor Tirgue va Calon Material)	8	0	8.12	<del>                                     </del>
	651101	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		+	5.12	+
1947	554101			+	<del> </del>	<del>                                     </del>
1		•		1	1	_L

Table 5

Table 5	5					
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1948	185432					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	26		26.65
1949	455598					
17.17	100070					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
1050	640254	10,17 (Colon Tullior Tissue Vs. Colon Mediatasis)		ТŤ		
1950	649354					<u> </u>
					6.24	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	ļ
1951	4408					ļ
						<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	17		4.36
		08,09 (Lung, High Met vs. Lung, Low Met)	21	3	9.78	<b>i</b> .
1952	452366					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6 44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
1953	452366	15,70 (Normal Colon Vo. Colon Tames 1,5000)				
1933	432300			-		<del> </del>
		15 17 Olymord Color Tienne ve Color Metectores	6	0	6.44	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	-
1954	727331			<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1955	644853					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1956	554079					1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	10		4.92
1957	556245	10,11 (001011111111111111111111111111111				
- 1,57	330213			_		t
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1050	557200	13,10 (Normal Colon vs. Colon Tuntor Tissue)	<del>-   °</del>	Ů		3.00
1958	557388					<del>                                     </del>
	<del> </del>	16 17 (Color Turner Transports Color Manatoria)	12	1 2	6.6	+
	ļ	16,17 (Colon Turnor Tissue vs. Colon Metastasis)	13	2	6.6	12.2
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	13		12.3
1959	449468			<b>_</b>		-
						ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
1960	556245					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1961	455327					
	1 2 2 2 7					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	12		11.18
		15,16 (Normal Colon Vs. Colon Tumor Tissue)	1	10		9.46
10/0	546622	15,10 (NOTHIAI COION VS. COION TURIOT TISSUE)		10	-	7.40
1962	546632					<del>                                     </del>
	<b></b>			-		1
	1	23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	3	34		11.22

Table 5

Table 5						1 75/4
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	15	0	15.23	
1963	558762					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
1964	550818	.0,11 (00.00				
1204	330010					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		Ť		7,00
1965	554079					-
						100
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	10		4.92
1966	452430					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	1
1967	452430					
1707						
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
1000	556000	13,17 (Normal Colon Tissue vs. Colon Metastasis)	<del>- + -</del>	Ť	0	
1968	556082					1
				10		0.05
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	10		9.85
1969	514418					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	. 9		8.86
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.38
1970	426895					
1570	420075					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	18		8.38
		13,17 (Normal Colon Tissue vs. Colon Mictastasis)	<del> </del> -			1
1971	560803					<del> </del>
						( 00
		16,17 (Colon Tumor Tissue vs Colon Metastasis)	0	7		6.89
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1972	447737					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	6	3.35	
1973	373432					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	23	78		3.16
		15,16 (Normal Colon vs. Colon Tumor Tissue)	23	53		2.18
			3	49		11.69
	990	08,09 (Lung, High Met vs. Lung, Low Met)	<del></del>	<del>- '</del>		11107
1974	779					<del> </del>
					2.66	-
		03,04 (Breast, High Met vs. Breast, Non-Met)	60	22	2.66	<del></del>
		01,02 (Colon, High Met vs. Colon, Low Met)	27	54		2.17
1975	455327			<u> </u>		
				<u>.</u>		<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10		9.46
		15.17 (Normal Colon Tissue vs. Colon Metastasis)	1	12		11.18
1976	554742					
17/0	334142			$\vdash$	<u> </u>	
		16.17 (Calam Tumon Tisque vs. Calam Matastassa)	0	6	<del>                                     </del>	5.91
		16,17 (Colon Tumor Tissue vs. Colon Metastasıs)	<del>-                                    </del>	<del>                                     </del>		1 3.71
1977	455327			-	<del> </del>	<del>                                     </del>
	L			-	<u> </u>	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10	<u> </u>	9.46
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	12		11.18
			1			

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLOX	227.41.5 2 41.5 1.7				
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
1979	727447	101,02 (Colon, High Met vs. Colon, 20w Met)				
17/7	727447	-				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1980	552905	10,17 (Colon Tulliol Tissue vs. Colon Metastasis)		H	0.07	
1960	332903					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
1001	446900	13,17 (NOTHIAI COIGH TISSUE VS. COIGH MICLASIASIS)	°	<del>- '-</del>		0.52
1981	440900					
		16 17 (Color Turon Tierra ve Color Motostoria)	2	11		5.42
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)				10.25
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	11		10.23
1982	644190					
					6.00	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
1983	455327					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10		9.46
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	12		11.18
1984	422375					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
1985	422375					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
1986	530774					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	5	16		2.98
1987	554101					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
1988	5268					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	15		16.26
1989	642461				,	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	14	0	15.03	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.16	
1990	770					<del>                                     </del>
1,,,,	,,,,					
		03,04 (Breast, High Met vs. Breast, Non-Met)	47	9	5.1	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	1	11.81	
1991	3837	13,17 (Normal Colon Fissue Vs. Colon Metastasis)		<u> </u>	11.01	
1991	3637					$\vdash$
		02 04 (Broost High Motars Broost Non Mot)	9	1	8.78	
1002	5(1202	03,04 (Breast, High Met vs. Breast, Non-Met)	-   '	1	0.76	
1992	561382					
		151/01		Ļ	( 2 *	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	2	6.34	<u> </u>
1993	4408					
				$\sqcup$		ļ
		08,09 (Lung, High Met vs. Lung, Low Met)	21	3	9.78	
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	17		4.36
1994	5686	1		1 1		1

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLSI	Libiar y 1 arr 11,22				
		03,04 (Breast, High Met vs. Breast, Non-Met)	16	3	5.2	
1005	374609	03,04 (Bleast, High Wet vs. Bleast, Non-Met)				
1995	374009					
		21,22 (Normal Prostate vs. Prostate Cancer)	1	9		9.15
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
1006	734793	15,16 (Normal Colon vs. Colon Turnor Tissue)		- '		
1996	734793					-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
1007	452420	13,10 (Normal Colon vs. Colon Tumor Hasue)		-		
1997	452430					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
	450040	15,17 (Normal Colon Tissue vs. Colon Metastasis)		Гď	0.11	
1998	450940					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.86
		15.17 (Colon Tumor Fissue vs. Colon Metastasis)	1	9		8.38
1000	460445	13,17 (Normal Coloil Fissue vs. Coloil Metastasis)				0.50
1999	460445					
		16.17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2000	540041	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		l ů		
2000	549041					-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2001	555376	16,17 (Colon Tullor Fissue vs. Colon Metastasis)		۲		
2001	555276					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2002	426895	10,17 (Colon Tullior Tissue vs. Colon Metastasis)		Ť		
2002	420893					<b>-</b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	18		8.38
2003	1833	13,17 (Normal Colon 1138de vs. Colon Medicales)				
2003	1833					
		01,02 (Colon, High Met vs. Colon, Low Met)	9	25		3.01
2004	446450	01,02 (Colon, 11igh Met vs. Colon, Low Met,				
2004	440430					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3.88	
2005	650517	15,10 (Normal Colon to Colon Tamor 13514)				
2003	030317					<u> </u>
	1	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
2006	554785	15,10 (11011mil Goldi 15) Goldi 151				
2000	334703					1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
<del></del>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
2007	607430	Total (Trontal Color Total				
2001	007.50					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
2008	446673	1-9, (				
2000	1	<del>                                     </del>		1		1
	-	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	1	8.12	1
2009	734685	10,17 (COIOII TUITOI TIBBUS 13. COIOII FICUADUS)				<b>†</b>
2009	134063			$t^-$		1
	-	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
	1	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	1
2010	11630	10,17 (Colon Tullor 11330C 13. Colon Housiasis)		<del>  </del>		<b> </b>

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		21,22 (Normal Prostate vs. Prostate Cancer)	1	12		12.2
		03,04 (Breast, High Met vs. Breast, Non-Met)	3	13		4.44
2011	2930					
		01,02 (Colon, High Met vs. Colon, Low Met)	3	14		5.06
2012	44424					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	41	90		2.16
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	25	90		3.35
		08,09 (Lung, High Met vs. Lung, Low Met)	4	40		7.16
2013	452052				***	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	1	8.59	
2014	449356					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	21		2.79
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	4	21		5.17
2015	726225					
					6.00	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	5.69
2016	152500	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
2016	453708			-		ļ
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	19	· · ·	18.71
		15,17 (Colon Turnor Tissue vs. Colon Metastasis)	1	19		17.7
2017	447858	13,17 (Normal Colon Tissue vs. Colon Metastasis)		17		17.7
2017	447030					<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	
2018	451613	15,70 (TOTAL COLOR TO COLOR TOTAL COLOR TO				
2010	.01015					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	14		3.26
2019	650337					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2020	62016					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2021	447250					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2022	3837					
				-	0.70	
2022	640614	03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
2023	640614					-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	0	13.2	
2024	729531	10,17 (Colon Tumor Tissue vs. Colon Metastasts)	13		13.4	
2024	147331			$\vdash \vdash$		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	5.00
2025	729531	20,2. (Colon Admor Alloue 101 Colon Plembully)		H	,	
20						<del>                                     </del>

Table 5

Table 5 SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLSI		0	6	AUD	5.68
		15,16 (Normal Colon vs. Colon Tumor Tissue)		0	6.09	3.08
2026	(47052	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	0.09	
2026	647952					
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
2027	446012	13,16 (Normal Cololl Vs. Cololl Tullot Tissue)		-	0.54	
2027	446913					
	<u> </u>	15,16 (Normal Colon vs. Colon Tumor Tissue)	63	1	66.59	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	10	0	8.55	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	10	0	11.43	<del> </del>
			63	0	67.62	
2029	2675	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0.5	<u> </u>	07.02	
2028	2675					
		102 04 (Parast Wall Makes Preset New Med)	12	2	5.85	
2020	642401	03,04 (Breast, High Met vs. Breast, Non-Met)	12		3.83	
2029	643481					
		Is is at the second sec		_	6.14	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2030	1345					
			10		2.05	
		21,22 (Normal Prostate vs. Prostate Cancer)	18	6	2.95	5.40
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	2.20	5 68
		08,09 (Lung, High Met vs. Lung, Low Met)	44	27	2.28	
		03,04 (Breast, High Met vs. Breast, Non-Met)	25	11	2.22	
2031	26					
					(0.10	
		03,04 (Breast, High Met vs. Breast, Non-Met)	62	0	60.49	20.25
		01,02 (Colon, High Met vs. Colon, Low Met)	0	28		30.36
2032	945					
			10	21		2.20
		01,02 (Colon, High Met vs. Colon, Low Met)	10	21		2.28
2033	449169					
		151601 IGIL GIL T	16	<u> </u>	0.46	
2024	204102	15,16 (Normal Colon vs. Colon Tumor Tissue)	16	2	8.46	
2034	394193					
		16 17 (O.1. 77 T)				5.01
2025	450010	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	<u> </u>	5.91
2035	452212			-		
	ļ	15 16 Olympia Colon un Colon Tyman Tigwel	6	0	6.34	
2026	204102	15,16 (Normal Colon vs. Colon Tumor Tissue)		ľ	0.54	-
2036	394193					
		16 17 (C. L. T Times of Color Metabolis)	0	6		5.91
2027	1210	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		0		3.91
2037	1310					
		02 04 (Durant High Maters Durant New Mat)	42	16	2.56	
		03,04 (Breast, High Met vs. Breast, Non-Met)				<b></b>
2022	724004	21,22 (Normal Prostate vs. Prostate Cancer)	15	2	7.38	-
2038	734094					-
		16 17 (Color Torres Tierres Color Maria		_	0.13	<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	<u> </u>
	1	23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	6	0	6.06	<u> </u>
		116 16 Olimont Colon on Colon William W.				
2039	646579	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57

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			<u> </u>	A (ID)	B/A
CLST					JD/A
	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
4471					1
	03,04 (Breast, High Met vs. Breast, Non-Met)	10	1	9.76	
729173					-
	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
		0	8		7.57
450323					<u> </u>
	15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
4652					<u> </u>
	03.04 (Breast, High Met vs. Breast, Non-Met)	16	2	7.81	
553316					<u> </u>
	15.16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	<u> </u>
642604					
	15.17 (Normal Colon Tissue vs. Colon Metastasis)	10	0	10.73	1
		13	0	13.2	
553316	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	10	13.2	╀
	4471 729173 450323 4652 553316	CLST Library Pair A,B  15,17 (Normal Colon Tissue vs. Colon Metastasis)  4471  03,04 (Breast, High Met vs. Breast, Non-Met)  729173  16,17 (Colon Tumor Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue)  450323  15,17 (Normal Colon Tissue vs. Colon Metastasis)  4652  03,04 (Breast, High Met vs. Breast, Non-Met)  553316  15,16 (Normal Colon vs. Colon Tumor Tissue)  642604  15,17 (Normal Colon Tissue vs. Colon Metastasis)  16,17 (Colon Tumor Tissue vs. Colon Metastasis)	CLST         Library Pair A,B         A           15,17 (Normal Colon Tissue vs. Colon Metastasis)         6           4471	CLST       Library Pair A,B       A       B         15,17 (Normal Colon Tissue vs. Colon Metastasis)       6       0         4471       ————————————————————————————————————	CLST         Library Pair A,B         A         B         A/B           15,17 (Normal Colon Tissue vs. Colon Metastasis)         6         0         6.44           4471

15,16 (Normal Colon vs. Colon Tumor Tissue)

03,04 (Breast, High Met vs. Breast, Non-Met)

15,17 (Normal Colon Tissue vs. Colon Metastasis)

15,16 (Normal Colon vs. Colon Tumor Tissue)

15,16 (Normal Colon vs. Colon Tumor Tissue)

15,17 (Normal Colon Tissue vs. Colon Metastasis)

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Table 5

Table 5		T T T A D	- I A I	В	A/B	B/A
SEQ	CLST	Library Pair A,B	A 2	11	A/D	5.64
		03,04 (Breast, High Met vs. Breast, Non-Met)		11		3.04
2056	452238					
		Late (a) I all a Cala Tarray	0	6		5.68
		15,16 (Normal Colon vs. Colon Tumor Tissue)	U			3.08
2057	1870					
		01 02 (C. L. Hick Maters Calon Love Mot)	11	31		3.06
		01,02 (Colon, High Met vs. Colon, Low Met)	12	3	3.9	3.00
20.50	550250	03,04 (Breast, High Met vs. Breast, Non-Met)	12		3.7	
2058	559259					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	17	5	3.59	
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	5	21		4.14
2059	453457	10,17 (Coloil Tullor Tissue vs. Coloil Metastasis)				
2039	433437					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
2060	8868	10,17 (Cololi Tunioi Tissue vs. Colon Metastasis)				
2060	8000					
		08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
2061	453059	03,04 (Bleast, High Met vs. Bleast, 1101 Fisch				
2001	433039					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
·-		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
2062	236368	13,10 (Normal Colon Isl Colon 12mol 12mol)				
2002	230300					
	_	18,19 (Normal Colon Tissue vs. Colon Tumor)	9	1	10.29	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	271	0	290.88	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	9	0	7.69	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	271	16	17.9	
	<del>                                     </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	16	0	16.25	
2063	453059					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	9	0	9.66	
2064	549979					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
2065	515631					
				<u> </u>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0_	6		5.68
2066	2235					
				ļ		
		03,04 (Breast, High Met vs. Breast, Non-Met)	36	12	2.93	
2067	448193					<u> </u>
				10		0.46
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	10	5.00	9.46
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	2	5.08	
2068	530774					
				1.0	<del>                                     </del>	3.00
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	5	16		2.98
2069	650204			<del> </del>		<del>                                     </del>
				<u></u>	11.01	<u> </u>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	0	11.81	<del> </del>
2070	644240			<u> </u>	L	<u> </u>

Table 5

Table 5	5					
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2071	552614					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	7		6.89
2072	727331					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
2073	185457					
				<u> </u>		
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	18		18.45
2074	454531					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
2075	643485					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	9	1	9.51	
		15,17 (Normal Colon Tissue vs Colon Metastasis)	9	0	9.66	
2076	733669					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2077	452344					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
2078	63602					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2079	454155					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	2	6.34	
2080	549903					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
2081	515631					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
2082	6878					
-		03,04 (Breast, High Met vs. Breast, Non-Met)	2	11		5.64
2083	2977					
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	4	3.66	
2084	553823					
	1			1		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	24	7	3.62	
	i	15,17 (Normal Colon Tissue vs. Colon Metastasis)	24	6	4.29	
2085	3070					
ļ				<u> </u>		
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2086	728884					
<u> </u>						
	<b> </b>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
	I	1 ' ` `			٠	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
2087	8166					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2088	644190					
						<u>.                                    </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	ļ
2089	733669			-		
		15 16 OL C. L. T T.		<del>  _</del>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	6.00	5.68
2090	728273	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	-
2090	120213			<b>!</b>		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.63
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	6.62
2091	406499	10,17 (Coloii Turioi Tissue vs. Coloii Miciasiasis)		<del>  '</del>	7.11	
2071	400422					<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<b></b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2092	557720			Ť	0.17	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2093	732050					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
-)-		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
2094	450867					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
2095	650297					
2006	440064	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2096	448064					
		15 1( Olympia Colored Colored Transport		<u> </u>	2.00	
2097	452530	15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3 88	
2097	432330					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	
2098	7592	15,10 (Ivotinal Colon vs. Colon Tuttor Tissue)	10	-	3.20	
2070	7072					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59
2099	733669					7.05
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2100	11028					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
2101	1013					
		01,02 (Colon, High Met vs. Colon, Low Met)	40	84		2.28
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2102	549265					

Table 5

Table 5 SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
DEQ	CLOX	220141 9 1444 1434				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3.88	
2103	376600	legate (Norman Colon to Colon Lands Tieste)				
2105	370000					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6 62
2104	643804	15,15 (Torrida Colon ISI Colon Tanta Tibeda)				
2104	043804		· · · · · · · · · · · · · · · · · · ·			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
	<u>                                     </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2105	454927	13,17 (Normal Colon 1135de vs. Colon Welastasis)		Ť	0111	
2103	757727					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	14		4.6
2106	446528	10,17 (Cololi Tutto) Tissue vs. Cololi Metastasis)		17		7.0
2106	440328					<u> </u>
		15 16 (Normal Colon va Colon Turnor Tigue)	10	0	10.57	1
2107	2210	15,16 (Normal Colon vs. Colon Tumor Tissue)	10	-	10.57	
2107	2218					
		O1 O2 (C-1 Ui-l- Max va C-1 I Max)	2	21		11.38
		01,02 (Colon, High Met vs. Colon, Low Met)			6.00	11.36
		08,09 (Lung, High Met vs. Lung, Low Met)	5	8	6.99	<u> </u>
	150504	03,04 (Breast, High Met vs. Breast, Non-Met)	27	8	3.29	<u> </u>
2108	452704			<u> </u>		<u> </u>
			<del></del>		10.57	ļ
	2.00.5	15,16 (Normal Colon vs. Colon Tumor Tissue)	10	1	10 57	<u> </u>
2109	84895					
				120		2.57
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	46	120		2.57
		08,09 (Lung, High Met vs. Lung, Low Met)	0	12		8 59
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	24	120		4.66
2110	157629					<u> </u>
				10		
		03,04 (Breast, High Met vs. Breast, Non-Met)	3	18		6.15
2111	2930					
		01,02 (Colon, High Met vs. Colon, Low Met)	3	14		5.06
2112	7037					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	9		9.76
2113	559806			<u> </u>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2114	452076			<u> </u>		ļ
	ļ			<u> </u>	4	
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	11	1	11.11	<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	20	7	2.9	
2115	454869			$\square$		ļ
	ļ					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
2116	559674					<u> </u>
				<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
2117	2235					
		03,04 (Breast, High Met vs. Breast, Non-Met)	36	12	2.93	L
2118	7545		1			l

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
	<u> </u>	01,02 (Colon, High Met vs. Colon, Low Met)	9	0	8.3	<del>                                     </del>
2119	729173					
						1
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	8	8.12	7.57
2120	650448	10,17 (Coloii Tullioi Tissue vs. Coloii Mcastasis)	8	Ť	6.12	<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<u> </u>
2121	172013			<u> </u>		ļ
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<u> </u>
2122	651088					
2122	(51000	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2123	651088					<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2124	726810					
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	7	7.11	6.62
2125	406499	10,17 (Colon Tullior House vs. Colon Hacustusis)		١Ů	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
2126	556325	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2120	330323					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
2127	644836					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
2128	649062					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	ļ
2129	454776	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	5	20		3.78
2130	377579					
	!	21,22 (Normal Prostate vs. Prostate Cancer)	25	53		2.16
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	13		6.06
2131	728131					
		15 16 (Normal Colon un Color Tresser Tiresse)				5.00
		15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	6	6.09	5.68
2132	475203	, ( )			2.07	
	-4-	15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	14		13 04
2133	727314	15,16 (Normal Colon vs. Colon Tumor Tissue)	1	12		11.35

	Ta	b	le	
ſ	CI		$\overline{}$	

Table 5 SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ.	CLSI	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
2134	552025					
213.	002020					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	6	3.35	
2135	561382					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	2	6.34	
2136	732579					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
2137	167					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	19		2.57
2138	185585					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2
2139	728131					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2140	475203					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	14		13.04
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	12		11.35
2141	724616					
				_	0.10	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	8		7.57
2143	645222					
			11		11.17	ļ
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0_	11.17	
2144	400362					
		16 17 (Color Towns Tissue us Color Matestagia)	35	117		3.29
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	21	117		5.19
2145	646592	15,17 (Normal Colon Tissue vs. Colon Metastasis)	21	117		3.17
2145	646583					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2146	475203	13,17 (Normal Colon 1135ac vs. Colon Weatsus)				
2140	473203			-		
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	14		13.04
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	12		11.35
2147	550001					
<u> </u>						
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	2	5.08	
2148	640703					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	0	10.16	
2149	646583					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2150	449468					<u> </u>

Table 5

Table 5		Yu D'AB	- I A	D	A/B	B/A
SEQ	CLST	Library Pair A,B	A	В	A/B	D/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
2151	449468	15,17 (Normal Coloir Fissue vs. Coloir Metastasis)				5.12
2131	77700					
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
2152	551628					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	38	5	8.03	ļ
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	38	13	3.14	ļ <u>.</u>
2153	449468					
				11		5 12
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
2154	417259					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	14		13.79
		15,17 (Colon Turnor Tissue vs. Colon Metastasis)	4	14		3.26
2155	448029	15,17 (Normal Colon Hissac vs. Colon Michaelas)				
2133	440023					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	10	1	10.16	
2156	524363					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2157	446531					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2158	561359			<b></b> -		
			12	1	6.34	<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.29	
2150	711297	15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	<del> </del>	7.27	
2159	/1129/					1
<b> </b>	ļ	15,16 (Normal Colon vs. Colon Tumor Tissue)	1	11		10.41
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
2160	650097					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2161	495715				<u> </u>	
<u> </u>		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
2162	734685			-		<del> </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.05	5.68
2163	560515	13,10 (Normal Colon vs. Colon Turnor Hasac)		Ť		
12103	300313					
	<b>†</b>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	-	5.91
2164	3441					
		03,04 (Breast, High Met vs. Breast, Non-Met)	8	0	7.81	
		01,02 (Colon, High Met vs. Colon, Low Met)	4	13		3.52
2165	729273			<b> </b>		<del> </del>
	ļ			<del>  _</del>	ļ	
<u> </u>	_	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7	7.11	6.62
1	L	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	7	0	7.11	<u> </u>

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
2166	557039					
2100	331037					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	16		7.88
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	2	5.81	
2167	711297	15,10 (Normal Colon vs. Colon Turnor Tissue)				
2167	711297					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	11		10.4
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
2160	711207	10,17 (Colon Turnor Tissue vs. Colon Metastasis)				<u> </u>
2168	711297					
		15 16 Olympia Colon vo Colon Turnor Tissue)	1	11		10.4
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	0	11 17	<u> </u>
24.60	2010	16,17 (Colon Tumor Tissue vs. Colon Metastasis)		Ť		┢┈┈
2169	2860					-
		Tiene)		13		12.3
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	13		12.1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	48		49.
		03,04 (Breast, High Met vs. Breast, Non-Met)	23	9	2.36	1 72.
		01,02 (Colon, High Met vs. Colon, Low Met)	23		2.50	<del>                                     </del>
2170	558534			-		$\vdash$
		1/ 17/Color Transa Times Color Materials	0	7		6.8
		16,17 (Colon Tumor Tissue vs. Colon Metastasıs)				1 0.0
2171	711297					-
		G1 M4 44 in	11	0	11.17	+-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	11	11.17	10.4
		15,16 (Normal Colon vs. Colon Tumor Tissue)		11		10.
2172	378457			-	-	+
		Ch. T. Time)	1	9		8.5
		15,16 (Normal Colon vs. Colon Tumor Tissue)		<del>  ´-</del>		1 0.0
2173	646583			-		+-
		Color Metators)	7	0	7.51	+-
	646700	15,17 (Normal Colon Tissue vs. Colon Metastasis)		<del>                                     </del>	7.51	+-
2174	646583			<u> </u>		
	<u> </u>	La sa Colon Metastaria	7	0	7.51	+
	1006	15,17 (Normal Colon Tissue vs. Colon Metastasis)		<del>  </del>	7.51	+
2175	1996			-		+
	<u> </u>	C.I. Matataia	13	4	3.3	<del>                                      </del>
	<u> </u>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	34	139	J.J	4.1
	<u> </u>	03,04 (Breast, High Met vs. Breast, Non-Met)	2	139		6.1
	<u> </u>	15,16 (Normal Colon vs. Colon Tumor Tissue)	20	13	2.15	1 0.1
		08,09 (Lung, High Met vs. Lung, Low Met)	20	113	2.13	<del> </del>
2176	7962			1-		+-
	<u> </u>	and the state of t	7	0	6.83	+-
		03,04 (Breast, High Met vs. Breast, Non-Met)	<del></del>	+ -	0.83	-
2177	645139			+		+
			<del></del>	-	0.14	1-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	9.14	+
2178	449468			+	<del>                                     </del>	+-
	<u> </u>		<del></del>	+	<del> </del>	1 .
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.1
2179	9898			+		+-
				1		+ -
		03,04 (Breast, High Met vs. Breast, Non-Met)	4	14		3.5
	406499	•				

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2181	1257					
		01,02 (Colon, High Met vs. Colon, Low Met)	46	20	2.12	
		21,22 (Normal Prostate vs. Prostate Cancer)	6	42		7.12
2182	549903					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
2183	557906					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	3	4.23	
2184	3538					ļ
		01,02 (Colon, High Met vs. Colon, Low Met)	18	5	3.32	ļ
2185	3114					
		03,04 (Breast, High Met vs. Breast, Non-Met)	9	1	8.78	
2186	426895					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	18		8.38
2187	923					
			<u> </u>			
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	23	2.6	3.11
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	23	9	2.6	<b> </b>
2188	645194			-		<del> </del>
		15 12 OL TO CL Marris		-	7.51	<del> </del>
	770161	15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2189	550161			-		
		15 16 Olympia Colonia Colonia Timora Timora	4	15		3.55
21.00	650110	15,16 (Normal Colon vs. Colon Tumor Tissue)	+	13		3.33
2190	650119			-		<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2191	642142	13,17 (Normal Colon Tissue vs. Colon Metastasis)		<del>ا</del>	7.31	
2191	042142			<del>                                     </del>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	l
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	11		5.2
2192	419255	15,10 (16.1.1.1 ed.en. to. ed.en. 14.1.1.1				
	113200					
	<u> </u>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	1	11.81	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	0	11.63	
2193	552905					
	f					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
2194	511997			Ľ		
	ĺ					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2195	551434					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	40	13	3.25	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	40	6	7.16	
2196	727447					

Table 5

Fable 5	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLSI	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
2197	378786	15,10 (Normal Colon Vs. Colon Tunior Tiesday)				
2197	318180					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.86
2198	649152	10,17 (Colon Tunio, Tiodae io Colon Tunio,				
2190	049132					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2100	18853	13,17 (Normal Colon Tissae vs. Colon Frenchis)				
2199	18833					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2200	642401	10,17 (Coloil Tulkoi Tissue vs. Coloil Metasusis)				
2200	643481			~ +		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
		15,17 (Normal Coloit Tissue vs. Coloit Wetastasis)		Ť		
2201	644417					
		16.15 (C.1. T Teaus vs. Colon Motostorys)	6	0	6.09	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)		Ť	0.03	
2202	726788					
		C. L. Matarasa	6	0	6.09	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6	0.03	5.68
		15,16 (Normal Colon vs. Colon Tumor Tissue)		<del>-                                    </del>		5.00
2203	206					
			19	4	6.64	
		08,09 (Lung, High Met vs. Lung, Low Met)	166	79	2.05	
		03,04 (Breast, High Met vs. Breast, Non-Met)	100	19	2.03	
2204	395930					
				0	6.44	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	<b>⊢</b> ° -	0.44	
2205	185589					<del> </del>
				9		9.22
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
2206	1441			_		
				10		4.02
		01,02 (Colon, High Met vs. Colon, Low Met)	9	40	2.22	4.82
		03,04 (Breast, High Met vs. Breast, Non-Met)	38	16	2.32	
2207	14522					
						-
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	8		8.2
2208	203605			ļ		
	<u> </u>				2.55	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	17	7	2.57	
2209	551527			<u></u>		<b>!</b>
				<u> </u>		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
2210	4509					
				ļ		
		08,09 (Lung, High Met vs. Lung, Low Met)	36	22	2.29	<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	40		41
2211	447737			<u> </u>		
						<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	19	6	3.35	
2212	447388					
<del></del>	1					
1						

Table 5

Table 5				-	4.00	75/1
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,16 (Normal Colon vs. Colon Tumor Tissue)	142	2	75.05	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	142	4	38.1	
		18,19 (Normal Colon Tissue vs. Colon Tumor)	18	0	20.57	
2213	451932					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	28	4	7.51	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	15	4	3.81	
2214	559043	10,17 (Colon Tullor Tissue vs. Colon Metastasio)		_	0.01	
2214	339043					
		(1) M(1)		50		4.49
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	50		4.48
		15,16 (Normal Colon vs. Colon Tumor Tissue)	54	11	5.19	ļ
2215	380634					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	14	_	4.6
2216	495715					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	8		7.88
2217	451932					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	28	4	7.51	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	15	4	3 81	
2210	447020	10,17 (Coloit Tuttioi Tissue vs. Coloit Wiciastasis)	15	-	3 01	
2218	447939					
		1516 AL 101 CI T T	1.4	<del>  ,</del>	140	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	14	1	14.8	7.00
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	1	8		7.88
2219	1181					
		03,04 (Breast, High Met vs. Breast, Non-Met)	17	58		3.5
2220	376600					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	7		6.62
2221	234761					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8 59	
2222	644417					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2223	639048					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7 51	<b> </b>
2224	11452	13,17 (Hollmar Colon Hissac vs. Colon Hicasasis)		Ť	, , ,	<u> </u>
2224	11432			<u> </u>		<u> </u>
		01 02 (Color High Maters Color Law Mat)	0	6		6.5
2225	450056	01,02 (Colon, High Met vs. Colon, Low Met)		0		0.5
2225	452076					<b>-</b>
			1.1	<del></del>	11 11	
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue)	11	1	11.11	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	20	7	2.9	<u> </u>
2226	644523					
				<u> </u>		
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2227	554678					
			-			Ι
	l	15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	31	5	6.05	
2229	450311	05,04 (Bleast, High Met vs. Bleast, 1401-1404)				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
2230	647280					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
2231	548858					
						40.0
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	11		10.83
2232	4204					-
		01,02 (Colon, High Met vs. Colon, Low Met)	16	2	7.38	
2233	540690					
						7.00
		23,24 (Normal Lung Tissue vs. Lung Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	8	11.17	7.92
2234	404774	10,17 (Colon Tumor Fissue vs. Colon Metastasis)				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	11	3	3.88	<u> </u>
2025	555000	15,17 (Normal Colon Tissue vs. Colon Metastasis)	11	1	11.81	
2235	557823					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	9		8.80
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	9		8.38
2236	1458					
		03,04 (Breast, High Met vs. Breast, Non-Met)	20	3	6.5	
2237	485431	03,0 1 (Bloady 11) Bit 110t 10t Breaty 1 of 10th				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	-	5.68
2238	2245					
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	
		01,02 (Colon, High Met vs. Colon, Low Met)	12	27		2.44
2239	3242					<u> </u>
	<u> </u>	01.02 (Colon Uick Met us Colon Low Met)	2	17		9.22
2240	648747	01,02 (Colon, High Met vs. Colon, Low Met)		11		7.22
2240	040747					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	ļ <u>.</u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<u> </u>
2241	3805			<b></b> -		-
	-	01,02 (Colon, High Met vs. Colon, Low Met)	17	3	5.23	<del>                                     </del>
2242	475203	2-7-2 (2000), 2000 (2000)				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	12		11.3
22.12	12010	15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	14	<u> </u>	13.0
2243	12018					<b> </b>
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.1
2244	475203					

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
SEQ	CLBI	2				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	12		11.35
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	14		13.04
2245	3805	15,17 (Normal Colon Tissue Vs. Colon Tissue Vs.				
2243	3003					
		01,02 (Colon, High Met vs. Colon, Low Met)	17	3	5.23	
2246	496132	or, oz (colon, riigii ivet is. colon, == s-s-y				
2240	470132					
	*	15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
2247	650600	13,10 (Normal colon 13, colon 14, co				
2241	030000					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2248	650749	13,17 (Normal Colon Fissac to Colon Fissac to				
2240	030749					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2249	223148	15,17 (Normal Colon Fissue 13. Colon Measures)				
2249	223146					1.00
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	9		8.38
2250	449	15,17 (Normal Colon Fissac Vis Colon Manually)				
2230	449					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	14		4.35
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	14		4.42
2251	735620	13,10 (Notifial Colon vs. Colon Tumor 115005)				
2231	733020					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6 09	
2252	650600	10,17 (Colon ramo) risodo var colon riconario,				1
LLJL	050000					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	
2253	218					
2233	210					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	11		10.25
		03,04 (Breast, High Met vs. Breast, Non-Met)	127	49	2.53	
2254	4161					
		08,09 (Lung, High Met vs. Lung, Low Met)	31	2	21.66	
		01,02 (Colon, High Met vs. Colon, Low Met)	13	1	11.99	
2255	373202					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	. 5	22		4.1
2256	724339					T
	├ <u></u>	15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2257	113291					
2207	1.02/1					
	1	15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	15		6.99
2258	736753					
	1					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	11		10.4
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	
2259	650600					
	320000					
	<del>                                     </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	

Table 5

Table 5				70	4 (TD)	TD/A
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
2260	451569					
	·					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0_	8		7.45
2261	1297					
		03,04 (Breast, High Met vs. Breast, Non-Met)	30	14	2.09	
2263	63602					
2203	05002					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	0	6		5.91
2264	2757	10,17 (Colon Tunior Tissue 18 Colon 18 Colon 18				
2204	2131					
		02 04 (Dreset High Met vs. Breest Non-Met)	4	16		4.1
22/2	272120	03,04 (Breast, High Met vs. Breast, Non-Met)	-	10		
2265	373128					
				20		5.19
		08,09 (Lung, High Met vs. Lung, Low Met)	4	29		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	14_		3.26
2266	641479					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	8	0	8.12	
2267	450380					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	31		7.22
		15,16 (Normal Colon vs. Colon Tumor Tissue)	4	23		5.44
2268	133512					
- 2200	133312					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	8		7
2260	447211	18,19 (Normal Colon Tissue vs. Colon Turkor)		-		
2269	447211					
		10 20 QL 1 Calar Tienus us Colon Motostogio)	9	0	7.69	
ļ		18,20 (Normal Colon Tissue vs. Colon Metastasis)	46	0	49.38	<del> </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	46	0	48.62	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)			10.29	<del> </del>
<u> </u>		18,19 (Normal Colon Tissue vs. Colon Tumor)	9	0	10.29	<del> </del>
2270	645222					<del> </del>
						<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	ļ
2271	645222					
						<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	11	0	11.17	<u> </u>
2272	17372					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2273	451619					
<u> </u>	<b>T</b>			Γ		
<del>                                     </del>		15,16 (Normal Colon vs. Colon Tumor Tissue)	24	9	2.82	
	<del>                                     </del>	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	9	39		4.27
2274	2510	Toyar (Colon Admin Alexandria)		<b>!</b>		
2214	2310			<del>                                     </del>		1
<b></b>	<del> </del>	02.04 (Deceat High Matics Deceat New Mat)	13	1	12.68	1
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	<del>                                     </del>	12.00	<del>                                     </del>
2275	643974			+-	ļ	<del> </del>
				<del>  _</del>		1
		15,16 (Normal Colon vs. Colon Turnor Tissue)	16	3	5.64	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	16	0	17.17	-
2276	500630			<u> </u>		<u> </u>

2277	3101 446938 554469	16,17 (Colon Tumor Tissue vs. Colon Metastasis)  01,02 (Colon, High Met vs. Colon, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue)	8 8 24 24	19 1 3 8	8.46 8.59 3.17	5.15
2278	446938 554469 554469	01,02 (Colon, High Met vs. Colon, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue)	8	19	8.46	5.15
2278	446938 554469 554469	01,02 (Colon, High Met vs. Colon, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8 59	5.15
2278 2 2279 5	446938 554469 554469	15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8 59	5.15
2279 5	554469 554469	15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8 59	5.15
2279 5	554469 554469	15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue)	24	3	8 59	
2279 5	554469 554469	15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	24	3	8 59	
2280 5	554469	15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	24	3	8 59	
2280 5	554469	15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)				
2280 5	554469	15,16 (Normal Colon vs. Colon Tumor Tissue)				
		15,16 (Normal Colon vs. Colon Tumor Tissue)				
		15,16 (Normal Colon vs. Colon Tumor Tissue)	24	8	3.17	
		15.17 (Normal Colon Tiesue ve Colon Matactacie)				1
2281		15 17 (Normal Colon Tissue vs. Colon Metastosis)	I			
2281		a de la la resolucia de Culon di Issauc vs. Colon iviciasiasis i	24	3	8.59	
2281		15,16 (Normal Colon vs. Colon Tumor Tissue)	24	8	3.17	
2201	2894	10,10 (110111111111111111111111111111111				
	2077					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	-	5.68
···		03,04 (Breast, High Met vs. Breast, Non-Met)	22	4	5.37	
		08,09 (Lung, High Met vs. Lung, Low Met)	5	20		2.86
2282	650600	Cung, Tigh Net vs. Eang, Ech Mety				
2282	030000					<b></b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	$\vdash$
2283	3101	13,17 (Normal Colon Tissue 18. Colon Medisalos)			****	
2203	3101					
		01,02 (Colon, High Met vs. Colon, Low Met)	4	19		5.15
2284	554469	01,02 (Cololi, High Met vs. Cololi, 20% Met)		1		
2207	334407					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	24	3	8.59	$\vdash$
		15,16 (Normal Colon vs. Colon Tumor Tissue)	24	8	3.17	
2285	9910	15,10 (trotting colon var colon rumer viscus)				
2203	<i>))</i> 10					┢
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	<u> </u>
2286	400608	os, or (Breast, Tright Free voi Breast, Treat	-			$\vdash$
2200	400000					$\vdash$
<del></del>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	21	7	3.22	$\vdash$
2287	555051	13,17 (101mar colon 1155de 15: colon intermental)				<del>                                     </del>
2207	333031					<b></b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	9		8.51
2288	185400	15,70 (Normal Colon 15, Colon 14, 16, 16, 16, 16, 16, 16, 16, 16, 16, 16				
2200	100-100			$\Box$		T
-		16.17 (Colon Tumor Tissue vs. Colon Metastasis)	1	9		8.86
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	62		63.5
2289	3059	05,04 (Breast, Figh Mot 18. Breast, From Mot)				
-207	2007			$\Box$		
		03,04 (Breast, High Met vs. Breast, Non-Met)	16	3	5.2	<del>                                     </del>
2290	647185	Oso ( (Stoudy High High to Stoudy Holl Haby	1.5	۱Ť		$\vdash$
2270	57/105			$\vdash$		<del>                                     </del>
<del>-  </del>		15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<del>                                     </del>
2291	1669	13,17 (troffisal Colon 1135uc vs. Colon (victastasis)		<u> </u>	3.77	<del> </del>
2271	1009			$\vdash$		<del>                                     </del>
$-\!+$		21,22 (Normal Prostate vs. Prostate Cancer)	0	34		34.5

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		03,04 (Breast, High Met vs. Breast, Non-Met)	40	0	39.03	
		01,02 (Colon, High Met vs. Colon, Low Met)	11	29		2.86
2292	7158					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
2293	496132					
				L		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	<u> </u>
2294	378623					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	33		10.25
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	24		7.57
2205	10.55	08,09 (Lung, High Met vs. Lung, Low Met)	1	16		11.45
2295	1257					<del>                                     </del>
		01 02 (Calan High Motors Calan Law Mot)	46	20	2.12	
		01,02 (Colon, High Met vs. Colon, Low Met)	6	20 42	2.12	7.12
2296	648499	21,22 (Normal Prostate vs. Prostate Cancer)	- 6	42		7.12
2290	046499					
		15,17 (Normal Colon Tissue vs. Colon Metastasıs)	10	0	10.73	-
		15,16 (Normal Colon vs. Colon Tumor Tissue)	10	2	5.28	<del>                                     </del>
2297	185627	15,10 (Avertical Colon Val. Colon Vallet Assac)		-	7.20	
	10002.					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
2298	640005					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	20	6	3.52	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	20	0	21.47	
2299	553462					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	6	0	6.34	
2300	649852	ļ				
		15.17.01			C 11	<b> </b>
2201	422275	15,17 (Normal Colon Tissue vs. Colon Metastasis)	6	0	6.44	<del> </del>
2301	422375					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	2	11		5.12
2302	10910	15,17 (Normal Colon Hissac Vs. Colon Measursis)				3.12
2502	10510					
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
2303	2737	1,02 (colon, 11,51, 11,01, 11,01, 11,01,01,01,01,01,01,01,01,01,01,01,01,0		H	<del></del>	-
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	3	14		4.6
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	4	14		3.26
2304	3438					
		01,02 (Colon, High Met vs. Colon, Low Met)	5	14		3.04
2305	3438					
		01,02 (Colon, High Met vs. Colon, Low Met)	5	14		3.04
2306	3763					
		01,02 (Colon, High Met vs. Colon, Low Met)	19	6	2.92	

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
2307	648966					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2308	724339					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
-		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2309	451569					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	8		7.45
2310	554109					1
		15,16 (Normal Colon vs. Colon Tumor Tissue)	15	2	7.93	
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	2	11		5.42
2311	380339					
						<u> </u>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	166	57	3.08	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	166	51	3.49	
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	9	0	6.73	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	18	0	15.39	
2312	729903	10,20 (Torrida Colon Tibble 18. Colon Medianes)		Ť	70.07	
2312	727703					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	<del> </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6	0.07	5.68
2313	45	15,10 (1101mai Colon vs. Colon Tumor Tissue)		۱Ů	-	3.00
2313	43			<del> </del>		
		08,09 (Lung, High Met vs. Lung, Low Met)	374	1067		2.04
		15,16 (Normal Colon vs. Colon Tumor Tissue)	40	119		2.04
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	40	209		4.87
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	29		25.38
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	0	11		12.87
		03,04 (Breast, High Met vs. Breast, Non-Met)	649	1876		2.96
2314	454653	03,04 (Bleast, High Met Vs. Bleast, Non-Met)	049	1870		2.90
2314	454055					-
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	3	17		5.28
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	5	17		3.35
2315	11526	10,17 (Coloii Turnoi Tissue vs. Coloii Wetastasis)		17		3.33
2313	11536					
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	<del> </del>
2316	373134	03,04 (Bleast, High Met Vs. Bleast, Non-Met)	- 0		3.63	<b></b>
2310	373134			_		<del>                                     </del>
		08,09 (Lung, High Met vs. Lung, Low Met)	2	45		16.1
			24	73		16.1
		15,17 (Normal Colon Tissue vs. Colon Metastasis)		$\vdash$		2.83
2217	105(01	16,17 (Colon Tumor Tissue vs. Colon Metastasis)	28	73		2.57
2317	185691					
		02.04 (P W. 1 M		Ļ		615
2210	00.17.1	03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		6.15
2318	234761		<u></u>			ļ
		16.15.07			0.50	ļ
22:5	<b></b>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	
2319	724339	<u> </u>				<u> </u>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
2320	732740					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
		16.17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2321	35895					
		21,22 (Normal Prostate vs. Prostate Cancer)	6	19		3.22
		15,16 (Normal Colon vs. Colon Tumor Tissue)	1	13		12.3
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	0	13.2	
2322	133512					
	100012					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	8		7
2323	2974	10,77 (Tormat colon 115500 151 colon 151111)				
2323	27,7					
		01,02 (Colon, High Met vs. Colon, Low Met)	22	7	2.9	
2324	500	01,02 (Colon, Tingh Met Vs. Colon, Low Met)		<u> </u>		
2324	300					<u> </u>
	l	03,04 (Breast, High Met vs. Breast, Non-Met)	89	22	3.95	
		01,02 (Colon, High Met vs. Colon, Low Met)	34	114	3.75	3.64
2325	376919	01,02 (Cololl, Tright Met vs. Cololl, Low Met)		111		3.01
2323	3/0919					<b></b> -
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	24	61		2.5
		21,22 (Normal Prostate vs. Prostate Cancer)	4	13		3.3
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	21	61		2.71
2226	8403	13,17 (Normal Colon Tissue vs. Colon Metastasis)	21	- O.		2.71
2326	8403					_
		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
2327	3643	03,04 (Bleast, High Met vs. Bleast, Non-Met)		<del>                                     </del>	3.03	
2321	3043					
		01,02 (Colon, High Met vs. Colon, Low Met)	6	19		3.43
2328	447211	01,02 (Colon, High Met Vs. Colon, Low Met)	<del>-   - °</del>	17		3.43
2328	447211			_		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	46	0	48.62	<del> </del>
	<del> </del>	15,17 (Normal Colon Tissue vs. Colon Metastasis)	46	0	49.38	<b>-</b>
	-	18,19 (Normal Colon Tissue vs. Colon Tumor)	9	0	10.29	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	9	0	7.69	
2329	447211	18,20 (Normal Colon Tissue vs. Colon Michaelasis)		l- Č	7.07	<u> </u>
2329	44/211			_		<del>                                     </del>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	46	0	49.38	<b></b>
		18,19 (Normal Colon Tissue vs. Colon Tumor)	9	0	10.29	
		18,20 (Normal Colon Tissue vs. Colon Metastasis)	9	0	7.69	
	-	15,16 (Normal Colon vs. Colon Tumor Tissue)	46	0	48.62	<del>                                     </del>
2220	14929	15,16 (Normal Colon vs. Colon Tulliol Tissue)	40	-	46.02	<b>-</b>
2330	14929			-		
		08,09 (Lung, High Met vs. Lung, Low Met)	23	16	2.01	<del>                                     </del>
			23	13	2.01	6.15
222.	(4003.1	15,16 (Normal Colon vs. Colon Tumor Tissue)	<del> </del>	13		0.13
2331	648934					<del> </del>
		15 16 QL 1 Q.L		^	0.47	<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	8	0	8.46	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	8	0	8.59	<del>                                     </del>
2332	731785					
				1		
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68

Table 5				~ T	4 (75)	75/4
SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	
2333	639908					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	13	4	3.44	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	13	0	13.95	
2334	344577					
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	28	2	14.22	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	38	2	20.39	
2335	2906					
2333	2700					
		01,02 (Colon, High Met vs. Colon, Low Met)	10	25		2.71
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
		03,04 (Breast, High Met vs. Breast, Non-Met)	2	15		7.69
2226	44(029	05,04 (Bleast, High Met vs. Bleast, Non Met)				
2336	446938					
		15.16 (Normal Colon vs. Colon Tumor Tissue)	8	1	8.46	
L	2.102	15,16 (Normal Colon vs. Colon Tullor Tissue)	<del></del>			
2337	2493					
		20 04 (D VII 1 M Durent New Med)	33	9	3.58	
		03,04 (Breast, High Met vs. Breast, Non-Met)	- 33		3.30	
2338	38					
			10	110		2.79
		15,16 (Normal Colon vs. Colon Tumor Tissue)	40	118		6.03
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	40	259		2.16
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	118	259	10.07	2.10
		19,20 (Colon Tumor Tissue vs. Colon Metastasis)	88	6	10.97	25.67
		18,19 (Normal Colon Tissue vs. Colon Tumor)	3	88		25.67
2339	13818			<u> </u>		
				_	5.05	
<u> </u>		03,04 (Breast, High Met vs. Breast, Non-Met)	6	0	5.85	
2340	8371			ļ		ļ
				ļ.,		
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6		6.5
2341	402494			<u> </u>		<u> </u>
						<del>                                     </del>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	3	13		4.1
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	13	3	4 4	<del> </del>
2342	731785			<del>-</del>		
				ļ.,		<del>                                     </del>
		16,17 (Colon Tumor Tissue vs. Colon Metastasis)	6	0	6.09	1 7.50
		15,16 (Normal Colon vs. Colon Tumor Tissue)	0	6		5.68
2343	4621			<u> </u>		ļ
				1-	<u> </u>	
		03,04 (Breast, High Met vs. Breast, Non-Met)	11	1	10.73	
2344	9750				ļ	<del></del>
		01,02 (Colon, High Met vs. Colon, Low Met)	0	6	<u> </u>	6.5
2345	133512			<del> </del>		<del>                                     </del>
					ļ	ļ
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	8		7
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
2346	162626				1	
		03,04 (Breast, High Met vs. Breast, Non-Met)	1	12	<u> </u>	12.3

15,16 (Normal Colon vs. Colon Tumor Tissue)   16,17 (Colon Tumor Tissue vs. Colon, Low Met)   08,09 (Lung, High Met vs. Breast, Non-Met)   03,04 (Breast, High Met vs. Colon, Low Met)   03,04 (Breast, High Met vs. Breast, Non-Met)   03,04 (Breast, High Met vs. Colon Metastasis)   15,16 (Normal Colon vs. Colon Tumor Tissue)   03,04 (Breast, High Met vs. Breast, Non-Met)   15,16 (Normal Colon vs. Colon Tumor Tissue)   03,04 (Breast, High Met vs. Breast, Non-Met)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,17 (Normal Colon vs. Colon Tumor Tissue)   15,17 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,17 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15	A	В	A/B	B/A
15,16 (Normal Colon vs. Colon Tumor Tissue) 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 2069  01,02 (Colon, High Met vs. Lung, Low Met) 08,09 (Lung, High Met vs. Breast, Non-Met) 2349 5868  03,04 (Breast, High Met vs. Breast, Non-Met) 2350 2683  01,02 (Colon, High Met vs. Breast, Non-Met) 2351 380409  16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2352 4389 15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon Tissue vs. Colon Metastasis) 2353 2353 2354 14929 15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Breast, Non-Met) 2355 134702 2356 642477 15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Breast, Non-Met) 2356 15,17 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 2357 14929 08,09 (Lung, High Met vs. Breast, Non-Met) 2358 134702 03,04 (Breast, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon Vs. Colon Tumor Tissue) 15,17 (Normal Colon Vs. Colon Tumor Tissue) 15,18 (Normal Colon Vs. Colon Tumor Tissue) 15,17 (Normal Colon Vs. Colon Tumor Tissue) 15,16 (Normal Colon Vs. Colon Tumor Tissue) 15,16 (Normal Colon Vs. Colon Tumor Tissue) 15,16 (Normal Colon Vs. Colon Tumor Tissue) 15,17 (Normal Colon Vs. Colon Tumor Tissue) 15,18 (Normal Colon Vs. Colon Tumor Tissue) 15,16 (Normal Colon Vs. Colon Tumor Tissue)				
16,17 (Colon Tumor Tissue vs. Colon Metastasis)  01,02 (Colon, High Met vs. Lung, Low Met) 08,09 (Lung, High Met vs. Lung, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met)  2349 5868  03,04 (Breast, High Met vs. Breast, Non-Met)  2350 2683  01,02 (Colon, High Met vs. Breast, Non-Met)  03,04 (Breast, High Met vs. Breast, Non-Met)  10,02 (Colon, High Met vs. Colon, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met)  15,16 (Normal Colon Tumor Tissue vs. Colon Metastasis) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Breast, Non-Met)  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 03,04 (Breast, High Met vs. Breast, Non-Met)				
16,17 (Colon Tumor Tissue vs. Colon Metastasis)  01,02 (Colon, High Met vs. Colon, Low Met) 08,09 (Lung, High Met vs. Breast, Non-Met) 03,04 (Breast, High Met vs. Breast, Non-Met)  2349 5868 03,04 (Breast, High Met vs. Breast, Non-Met)  2350 2683 01,02 (Colon, High Met vs. Breast, Non-Met) 03,04 (Breast, High Met vs. Breast, Non-Met)  2351 380409 16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)  2352 63991 15,17 (Normal Colon Tissue vs. Colon Metastasis) 2353 535 03,04 (Breast, High Met vs. Breast, Non-Met) 2354 14929 15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met) 2356 642477 15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Breast, Non-Met) 2357 14929 08,09 (Lung, High Met vs. Colon Tumor Tissue) 15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702 03,04 (Breast, High Met vs. Breast, Non-Met)	0	6		5.68
01,02 (Colon, High Met vs. Colon, Low Met)	6	0	6.09	
01,02 (Colon, High Met vs. Colon, Low Met)   08,09 (Lung, High Met vs. Lung, Low Met)   03,04 (Breast, High Met vs. Breast, Non-Met)   2349   5868   03,04 (Breast, High Met vs. Breast, Non-Met)   2350   2683   01,02 (Colon, High Met vs. Colon, Low Met)   03,04 (Breast, High Met vs. Breast, Non-Met)   2351   380409   16,17 (Colon Tumor Tissue vs. Colon Metastasis)   15,16 (Normal Colon vs. Colon Tumor Tissue)   2352   639991   15,17 (Normal Colon Tissue vs. Colon Metastasis)   2353   535   2354   14929   15,16 (Normal Colon vs. Colon Tumor Tissue)   2354   14929   2355   134702   2356   642477   2356   642477   2356   642477   2357   14929   2358   134702   2358   134702   2358   134702   2358   134702   2358   134702   2358   134702   2359   185649   2359   185649   2359   185649   2359   185649   2350   23,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649   2359   185649   2350   23,04 (Breast, High Met vs. Breast, Non-Met)				
08,09 (Lung, High Met vs. Lung, Low Met)				
08,09 (Lung, High Met vs. Lung, Low Met)	26	8	3	
03,04 (Breast, High Met vs. Breast, Non-Met)	21	6	4.89	
2349 5868  03,04 (Breast, High Met vs. Breast, Non-Met)  2350 2683  01,02 (Colon, High Met vs. Colon, Low Met)  03,04 (Breast, High Met vs. Breast, Non-Met)  2351 380409  16,17 (Colon Turnor Tissue vs. Colon Metastasis)  15,16 (Normal Colon vs. Colon Turnor Tissue)  2352 639991  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Turnor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Turnor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Breast, Non-Met)  15,16 (Normal Colon Tissue vs. Colon Metastasis)  2358 134702  03,04 (Breast, High Met vs. Lung, Low Met)  15,16 (Normal Colon Vs. Colon Turnor Tissue)  03,04 (Breast, High Met vs. Breast, Non-Met)	1	47	1,.07	48.17
03,04 (Breast, High Met vs. Breast, Non-Met)		47		40.17
2350 2683  01,02 (Colon, High Met vs. Colon, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met)  2351 380409  16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)  2352 639991  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon Tissue vs. Colon Metastasis)  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649				
2350 2683  01,02 (Colon, High Met vs. Colon, Low Met) 03,04 (Breast, High Met vs. Breast, Non-Met)  2351 380409  16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)  2352 639991  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon Tissue vs. Colon Metastasis)  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649	10	_	4.00	
01,02 (Colon, High Met vs. Colon, Low Met)   03,04 (Breast, High Met vs. Breast, Non-Met)   2351   380409   16,17 (Colon Tumor Tissue vs. Colon Metastasis)   15,16 (Normal Colon vs. Colon Tumor Tissue)   2352   639991   15,17 (Normal Colon Tissue vs. Colon Metastasis)   2353   535     03,04 (Breast, High Met vs. Breast, Non-Met)   2354   14929   15,16 (Normal Colon vs. Colon Tumor Tissue)   08,09 (Lung, High Met vs. Lung, Low Met)   2355   134702     03,04 (Breast, High Met vs. Breast, Non-Met)   2356   642477     15,16 (Normal Colon vs. Colon Tumor Tissue)   15,17 (Normal Colon Tissue vs. Colon Metastasis)   2357   14929     08,09 (Lung, High Met vs. Lung, Low Met)   15,16 (Normal Colon vs. Colon Tumor Tissue)   2358   134702     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350	10	2	4.88	
03,04 (Breast, High Met vs. Breast, Non-Met)				
03,04 (Breast, High Met vs. Breast, Non-Met)				
16,17 (Colon Tumor Tissue vs. Colon Metastasis)   15,16 (Normal Colon vs. Colon Tumor Tissue)   2352   639991     15,17 (Normal Colon Tissue vs. Colon Metastasis)   2353   535     03,04 (Breast, High Met vs. Breast, Non-Met)   2354   14929     15,16 (Normal Colon vs. Colon Tumor Tissue)   08,09 (Lung, High Met vs. Lung, Low Met)   2355   134702     134702     15,16 (Normal Colon vs. Colon Tumor Tissue)   15,16 (Normal Colon vs. Colon Tumor Tissue)   15,17 (Normal Colon Tissue vs. Colon Metastasis)   2357   14929     15,16 (Normal Colon Tissue vs. Colon Metastasis)   2358   134702     03,04 (Breast, High Met vs. Lung, Low Met)   2358   134702     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     03,04 (Breast, High Met vs. Breast, Non-Met)   2359   185649     2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   2350   235	3	22		7.95
16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649	13	2	6.34	
16,17 (Colon Tumor Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702  03,04 (Breast, High Met vs. Lung, Low Met) 2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)				_
15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon Tissue vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649				<u> </u>
15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon Vs. Colon Tumor Tissue)  15,16 (Normal Colon Vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649	10	2	5.08	
2352 639991  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Turnor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Turnor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Turnor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Turnor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649	0	10		9.46
15,17 (Normal Colon Tissue vs. Colon Metastasis)  2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649				
2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649				
2353 535  03,04 (Breast, High Met vs. Breast, Non-Met)  2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649	6	0	6.44	
03,04 (Breast, High Met vs. Breast, Non-Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  03,04 (Breast, High Met vs. Breast, Non-Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649		-		ļ —
2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)		-		<del></del>
2354 14929  15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)	87	13	6.53	<del>                                     </del>
15,16 (Normal Colon vs. Colon Tumor Tissue) 08,09 (Lung, High Met vs. Lung, Low Met)  2355 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)	- 87	13	0.55	<del>                                     </del>
08,09 (Lung, High Met vs. Lung, Low Met)  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)		<del> </del>		<u> </u>
08,09 (Lung, High Met vs. Lung, Low Met)  03,04 (Breast, High Met vs. Breast, Non-Met)  2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)		<b> </b>		(15
2355 134702 03,04 (Breast, High Met vs. Breast, Non-Met) 2356 642477 15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis) 2357 14929 08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702 03,04 (Breast, High Met vs. Breast, Non-Met) 2359 185649 03,04 (Breast, High Met vs. Breast, Non-Met)	2	13	ļ	6.15
03,04 (Breast, High Met vs. Breast, Non-Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)	23	16	2.01	ļ
2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)				
2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)				
2356 642477  15,16 (Normal Colon vs. Colon Tumor Tissue)  15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		7.17
15,16 (Normal Colon vs. Colon Tumor Tissue) 15,17 (Normal Colon Tissue vs. Colon Metastasis) 2357 14929  08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met) 2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)				
15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)				
15,17 (Normal Colon Tissue vs. Colon Metastasis)  2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	7.4	
2357 14929  08,09 (Lung, High Met vs. Lung, Low Met)  15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	7.51	
08,09 (Lung, High Met vs. Lung, Low Met) 15,16 (Normal Colon vs. Colon Tumor Tissue) 2358 134702 03,04 (Breast, High Met vs. Breast, Non-Met) 2359 185649 03,04 (Breast, High Met vs. Breast, Non-Met)		+-		
15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)		<b>-</b>		
15,16 (Normal Colon vs. Colon Tumor Tissue)  2358 134702  03,04 (Breast, High Met vs. Breast, Non-Met)  2359 185649  03,04 (Breast, High Met vs. Breast, Non-Met)	23	16	2.01	1
2358 134702 03,04 (Breast, High Met vs. Breast, Non-Met) 2359 185649 03,04 (Breast, High Met vs. Breast, Non-Met)	2	13		6.15
03,04 (Breast, High Met vs. Breast, Non-Met) 2359 185649 03,04 (Breast, High Met vs. Breast, Non-Met)		15		+
2359 185649 03,04 (Breast, High Met vs. Breast, Non-Met)		+-		+
2359 185649 03,04 (Breast, High Met vs. Breast, Non-Met)		+-		7.1
03,04 (Breast, High Met vs. Breast, Non-Met)	0	7		/.1.
		<del> </del>		<del></del>
	0	7	<u> </u>	7.13
		<u>L</u>		<u> </u>
03,04 (Breast, High Met vs. Breast, Non-Met)	0	15		15.3
		1		$\top$
2361 643955		$\top$	<b>†</b>	$\top$
Accept 10.1 Till Old Manager	12	0	12.88	+
15,17 (Normal Colon Tissue vs. Colon Metastasis) 15,16 (Normal Colon vs. Colon Tumor Tissue)	12	1 0	12.68	+-

Table 5	;		
SEQ	CLST	Library Pair A,B	
2362	643955		
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	
		15 16 (Normal Colon ve. Colon Tumor Tienus)	

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
2362	643955					
						<b></b>
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	0	12.88	<b></b>
		15,16 (Normal Colon vs. Colon Tumor Tissue)	12	0	12.68	1
2363	4455					1
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	
2364	185567					
		03,04 (Breast, High Met vs. Breast, Non-Met)	0	9		9.22
2365	9115	<u> </u>				
		01,02 (Colon, High Met vs. Colon, Low Met)	0	7		7.59
2366	14929			<u> </u>		L
				<u> </u>		
		08,09 (Lung, High Met vs. Lung, Low Met)	23	16	2.01	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	13		6.15
2367	14929				<u> </u>	L
					ļ	
		08,09 (Lung, High Met vs. Lung, Low Met)	23	16	2.01	
		15,16 (Normal Colon vs. Colon Tumor Tissue)	2	13	<u> </u>	6.15
2368	4181	<del> </del>		<b> </b>	<u></u>	ļ
	<u> </u>				4.00	ļ
22(0	5207	03,04 (Breast, High Met vs. Breast, Non-Met)	15	3	4.88	
2369	5206	<del> </del>				<del> </del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	<del>                                     </del>
2370	825	03,04 (Bleast, High Wet vs. Breast, Non-Wet)		-	0.83	
2510	023			<u> </u>		<del> </del>
		03,04 (Breast, High Met vs. Breast, Non-Met)	70	25	2.73	<b></b> -
		08,09 (Lung, High Met vs. Lung, Low Met)	13	3	6.06	<del> </del>
2371	825	3,3		<u> </u>		
		<u> </u>				
		08,09 (Lung, High Met vs. Lung, Low Met)	13	3	6.06	
		03,04 (Breast, High Met vs. Breast, Non-Met)	70	25	2.73	
2372	825					
		03,04 (Breast, High Met vs. Breast, Non-Met)	70	25	2.73	
		08,09 (Lung, High Met vs. Lung, Low Met)	13	3	6.06	
2373	2748					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	26		14.09
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	4	3.66	
2374	2748					
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	4	3.66	
		01,02 (Colon, High Met vs. Colon, Low Met)	2	26		14.09
2375	2748					
		01,02 (Colon, High Met vs. Colon, Low Met)	2	26		14.09
2376	122512	03,04 (Breast, High Met vs. Breast, Non-Met)	15	4	3.66	
23/0	133512	<del> </del>				
		18,19 (Normal Colon Tissue vs. Colon Tumor)	0	8		7
		120512 (13011161 Colon 11880c vs. Colon 10110r)	1 0	0		

Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	0	7		6.52
2377	2748					
2377	27-10					
		03,04 (Breast, High Met vs. Breast, Non-Met)	15	4	3.66	-
			2	26	3.00	14.00
		01,02 (Colon, High Met vs. Colon, Low Met)		20		14.09
2378	642477					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	
2379	642477					
		15,16 (Normal Colon vs. Colon Tumor Tissue)	7	0	7.4	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	7	0	7.51	<u> </u>
2200	2402	13,17 (Normal Colon Fissuc vs. Colon Miciasiasis)		Ů	7.51	
2380	2493		· · · · · · · · · · · · · · · · · · ·			
				_		
		03,04 (Breast, High Met vs. Breast, Non-Met)	33	9	3.58	ļ
2381	5796					
		08,09 (Lung, High Met vs. Lung, Low Met)	5	0	6.99	
		03,04 (Breast, High Met vs. Breast, Non-Met)	14	3	4.55	
2382	3782					<u> </u>
2302	3702					<del>                                     </del>
-		02.04 (D			20.40	-
		03,04 (Breast, High Met vs. Breast, Non-Met)	21	0	20.49	
2383	884					
		03,04 (Breast, High Met vs. Breast, Non-Met)	51	20	2.49	
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	1	10		9.32
		21,22 (Normal Prostate vs. Prostate Cancer)	20	7	2.81	
		08,09 (Lung, High Met vs. Lung, Low Met)	10	63		4.51
2384	5860	(				t
230-7	3000					<del></del>
		02.04 (December Health Markets December Mark)	10	2	4.88	
		03,04 (Breast, High Met vs. Breast, Non-Met)	10		4.00	
2385	5275					ļ
		01,02 (Colon, High Met vs. Colon, Low Met)	11	2	5.07	
2386	3932					
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	
2387	884					
		08,09 (Lung, High Met vs. Lung, Low Met)	10	63		4.51
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	10	10		9.32
				-	2.40	7.52
		03,04 (Breast, High Met vs. Breast, Non-Met)	51	20	2.49	<u> </u>
		21,22 (Normal Prostate vs. Prostate Cancer)	20	7	2.81	
2388	4455					
		03,04 (Breast, High Met vs. Breast, Non-Met)	13	1	12.68	
2389	5860					
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	2	4.88	
2200	50(0	OF (Dicast, High refer vs. Dicast, Holl-refer)	10	┝╧┤	7.00	<b> </b>
2390	5860			$\vdash$		<u> </u>
				$\vdash$		<u> </u>
		03,04 (Breast, High Met vs. Breast, Non-Met)	10	2	4.88	<u> </u>
2391	372791					

## Table 5

SEQ	CLST	Library Pair A,B	A	В	A/B	B/A
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	0	12.88	
2392	5206					
-		03,04 (Breast, High Met vs. Breast, Non-Met)	7	0	6.83	
2393	372791					
		15,17 (Normal Colon Tissue vs. Colon Metastasis)	12	0	12.88	
2394	2846					
<del></del>		01,02 (Colon, High Met vs. Colon, Low Met)	8	19		2.57
2395	5275					
		01,02 (Colon, High Met vs. Colon, Low Met)	11	2	5 07	
2396	2846					
		01,02 (Colon, High Met vs. Colon, Low Met)	8	19		2 57